SPECIFICATIONS AND FABRICATION PROCEDURES FOR SM-1A CORE II NEUTRON ABSORBER SECTIONS

Materials Technology Branch
Nuclear Power Engineering Department

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Post Office Box 414
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9
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Washington 25, D. C.

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Union Carbide Nuclear Corporation
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P. O. Box "Y"
Oak Ridge, Tennessee
Attention: A. L. Boch

14
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Technical Information Services Extension
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17-18
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Attention: Mr. G. E. Richards
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Albany 1, New York

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FOREWORD

These specifications were prepared to aid in the procurement of the Control Rod Absorber Section for Core II of SM-1A from an industrial manufacturer.

The material contained herein is based on ORNL-2733 and incorporates changes which are the result of design and material changes in addition to process changes resulting from experience gained during the manufacture of similar components by a commercial fabricator.
ABSTRACT

The control rod absorber section consists of composite plates joined by welding to form a rectangular parallelepiped. The composite plates consist of compacts of europium oxide in a stainless steel matrix that are clad with stainless steel by hot roll-bonding.

These specifications cover materials and process required to produce the absorber sections. The procedural specifications are specific for manufacturing these components, and represent a detailed guide for fabricators. It is recognized that because of differences in equipment, fabricators may be required to modify some of the detailed procedures to arrive at the same result. Prior approval for any deviation from these specifications must be approved by the Contracting Agency, or its authorized representative.
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# Absorber Section Specifications

## A. Core Material Requirements

1. **Neutron Absorber**
   - Europium content per plate, g: 151.00
   - Allowable tolerances:
     - Weighing: ± 0.3%
     - Eu/europium oxide: ± 0.5%
     - Handling losses: - 0.1%

2. **Matrix Material**
   - Stainless steel powder (prepared from elementals) makeup, per plate:
     - Iron, g: 220.95
     - Chromium, g: 56.01
     - Nickel, g: 34.23
   - Allowable tolerances:
     - Weighing: ± 0.9%
     - Handling loss: - 0.3%

## B. Material Specifications

1. **Neutron Absorber**
   - The neutron absorbing material shall be a thorium-free high-purity europium oxide containing a nominal 98 wt. % europium oxide. Received europium oxide, "dead-burned" from the oxalate, shall be high fired in a tungsten crucible under a dry hydrogen atmosphere for 3 hours at 3090°F. After firing, the material shall be comminuted to a particle size of less than 44 \( \mu \) by hammer forging and intermediate screening to avoid excessive fines. This material should contain a minimum 85 wt. % europium. The thorium content is determined by alpha activity measurements and material with alpha counts in excess of background is considered contaminated and questionable.

2. **Matrix Material**
   - The matrix material and diffusion barrier foil material shall be prepared from elemental iron, chromium and nickel powders in the weight ratio of 71, 18 and 11, respectively. The particle size of all materials is -325 mesh. The iron powder should be hydrogen annealed electrolytic iron with a mini-

* Allowable tolerances were established by ORNL as a result of uncertainties involved in fabricating absorber sections for Core II of SM-1.
imum iron content of 99.0 W/o. The nickel powder shall be prepared by the carbonyl process and the chromium powder may be produced either electrolytically or by lithium reduction. The maximum content of impurities for each element is as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Desired Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silicon</td>
<td>0.03 w/o</td>
</tr>
<tr>
<td>Oxygen</td>
<td>0.10 w/o</td>
</tr>
<tr>
<td>Carbon</td>
<td>0.02 w/o</td>
</tr>
<tr>
<td>Cadmium</td>
<td>0.002 w/o</td>
</tr>
<tr>
<td>Boron</td>
<td>0.002 w/o</td>
</tr>
<tr>
<td>Rare Earths</td>
<td>0.002 w/o</td>
</tr>
</tbody>
</table>

3. **Diffusion Barrier Foil**

The foil is prepared by blending, pressing, sintering, rolling, and annealing the above matrix material. The foil is rolled to a thickness of 0.005 "± 0.0005".

4. **Wrought Stainless Steel**

This specification covers a high quality stainless steel to be furnished in the form of sheared mill plate or sheet.

This steel is AISI Type 347 having the chemistry limits set forth in ASTM A-240-58T except that the cobalt shall be 0.025 w/o max. and tantalum shall be 0.01 w/o max.

**Chemical Composition: - in w/o**

<table>
<thead>
<tr>
<th>Element</th>
<th>ASTM Specification</th>
<th>Desired Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>0.08 max.</td>
<td>0.06 max.</td>
</tr>
<tr>
<td>Nickel</td>
<td>9.00-13.00</td>
<td>9.25-9.75</td>
</tr>
<tr>
<td>Chromium</td>
<td>17.00-19.00</td>
<td>18.00-18.75</td>
</tr>
<tr>
<td>Manganese</td>
<td>2.0 max.</td>
<td>1.00-1.50</td>
</tr>
<tr>
<td>Phosphorous</td>
<td>0.045 max.</td>
<td>0.030 max</td>
</tr>
<tr>
<td>Sulfur</td>
<td>0.030 max.</td>
<td>0.030 max</td>
</tr>
<tr>
<td>Silicon</td>
<td>1.0 max.</td>
<td>0.50-0.80</td>
</tr>
<tr>
<td>Columbium + Tantalum</td>
<td>10 x C min; 1.10 max.</td>
<td>0.60-0.80</td>
</tr>
<tr>
<td>Cobalt</td>
<td></td>
<td>0.025 max.</td>
</tr>
<tr>
<td>Tantalum</td>
<td></td>
<td>0.04 max.</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>0.50 max.</td>
<td>0.20 max.</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>0.50 max.</td>
<td>0.20 max.</td>
</tr>
<tr>
<td>Copper</td>
<td>0.50 max.</td>
<td>0.20 max.</td>
</tr>
</tbody>
</table>

The material with the above composition limits will be used for inert gas shielded tungsten arc welding without filler metal. In addition to meeting the chemical restrictions set forth in this specification, the delta ferrite content of this material, as calculated from the Schaeffler Constitution Diagram shall be a minimum of 2% and a maximum of 10%.
In calculating the delta ferrite content from the Schaeffler Constitution Diagram, the following multiplying factors shall be used in computing the chromium and nickel equivalents for the production order:-

<table>
<thead>
<tr>
<th>Element</th>
<th>Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>30</td>
</tr>
<tr>
<td>Manganese</td>
<td>1/2</td>
</tr>
<tr>
<td>Silicon</td>
<td>1-1/2</td>
</tr>
<tr>
<td>Chromium</td>
<td>1</td>
</tr>
<tr>
<td>Nickel</td>
<td>1</td>
</tr>
<tr>
<td>Cb + Ta</td>
<td>1/2</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>30</td>
</tr>
<tr>
<td>Copper</td>
<td>2</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>1</td>
</tr>
</tbody>
</table>

In the event that the chemistry of the heat is outside the range of the desired specification, but within the ASTM chemistry, and contains a delta ferrite content in the range of 2 to 10% as calculated from the Schaeffler Diagram, the heat will be acceptable.

In the event the chemistry is outside the desired specification limits, and below the lower limit of 2% delta ferrite, a weldability test shall be made without filler addition. Freedom from cracking shall consider the heat acceptable. The type and details of the welding test shall be as mutually agreed upon between Allegheny Ludlum and Alco Products, Inc., Material that is not weldable, as determined by this test will not be acceptable.

Inclusion Content:-

The inclusion content of this material in the form of 1" thick sheet bar as determined by Method "B" set forth in ASTM E-45-51 shall be equal to or better than an inclusion rating of (12-512). This rating as determined by method 'B' shall be comparable to the following method "A" rating:-

<table>
<thead>
<tr>
<th>Inclusion Type</th>
<th>Thin Series</th>
<th>Heavy Series</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type A (Sulfide)</td>
<td>2</td>
<td>1-1/2</td>
</tr>
<tr>
<td>Type B (Alumina)</td>
<td>2-1/2</td>
<td>2</td>
</tr>
<tr>
<td>Type C (Silicate)</td>
<td>3</td>
<td>3-1/2</td>
</tr>
<tr>
<td>Type D (Globular Oxide)</td>
<td>3</td>
<td>2</td>
</tr>
</tbody>
</table>
The thickness of the inclusions found in 1" thick sheet bar shall not be any greater than that specified in the Inclusion Chart of ASTM E-45-51 specification.

**Physical Properties:**

<table>
<thead>
<tr>
<th>Material</th>
<th>Tensile Strength</th>
<th>0.2% Offset Yield Strength</th>
<th>% Elongation in 2&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plate, Sheet, and Strip</td>
<td>75,000 psi</td>
<td>30,000 psi</td>
<td>40%</td>
</tr>
</tbody>
</table>

**Purchase Condition:**

Cold rolled, annealed, and pickled sheared mill plate or sheet.

C. **Dimensional Requirements**

The component parts shall be manufactured and assembled in accordance with the dimensional specifications set forth on Alco Drawings listed below:

<table>
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<th>Description of Item</th>
<th>Reference Drawing</th>
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<tr>
<td>Absorber Plates</td>
<td>Alco Drawing D9-13-2017-C</td>
</tr>
<tr>
<td>Pin</td>
<td>Alco Drawing A9-13-2019-B</td>
</tr>
<tr>
<td>Absorber Section Assembly</td>
<td>Alco Drawing D9-13-1002-B</td>
</tr>
</tbody>
</table>

Since these drawings are subject to revision, it is advisable that potential fabricators contact the Contracting Agency, in order to obtain the latest prints before initiating any fabrication work.

D. **Finish Requirements**

All machined surfaces in contact with the coolant shall have a finish of at least 125 RMS except as noted in the drawings.
E. Qualification of Absorber Plate Fabrication Procedure

1. Introduction

Quality control of the absorber section is primarily attained by rigid adherence to proven fabrication procedures. Thus, specifications for the absorber sections cannot be prepared along the lines normally used for industrial products, in which quality is assured by nondestructive inspection to industry-wide standards. Consequently, the manufacturer will be required to qualify the fabrication procedure which is to be employed in production.

2. Method of Qualification

Qualification shall be performed by subjecting three sample plates to the tests outlined in 4-a through 4-i of this section. Approval of qualification shall be given by the Contracting Agency, or its authorized representative when it has been demonstrated to the satisfaction of the Contracting Agency, or its authorized representative that the requirements of 4-a through 4-i have been met.

3. Preparation of Qualification Samples

The three sample plates shall be prepared in conformance with the exact and complete fabrication procedure which is proposed to be used in the manufacture of the absorber sections. Europium oxide, identical in every respect with the material to be used in the absorber sections, shall be used in the sample plates. The sample plates shall be subjected to the identical high-temperature treatment which will be encountered in the proposed procedure for fabrication of the composite plate. Approval of qualification must be obtained prior to inception of the manufacture of absorber plates unless otherwise specifically authorized by the Contracting Agency, or its authorized representative. Approval by the Contracting Agency, or its authorized representative of the procedure as used in establishing qualification or of any modification thereof will not relieve the manufacturer of any responsibility for any phase of the fabrication of the absorber section or for conformity to specification requirements.

4. Tests for Qualification

a. Visual Inspection – The three sample plates shall be inspected for over-all width, length, and thickness dimensions and shall be inspected for pitting, surface condition, and finish. The three sample plates shall meet all dimensional requirements and shall not exhibit any oxide indentations or pits in excess of 0.005 in. deep, scratches over 0.002 in. deep, blisters, scale, or dents.
b. Radiographic Examination - The three sample plates shall then be radiographed to delineate the europium oxide-bearing core area and voids or other internal defects. The radiographs shall be made using proper techniques with a fine-grained film such as Eastman "M". The radiographs shall be used as the basis of measuring core length, core width, and inactive edges and ends.

The samples shall meet all dimensional requirements and shall exhibit no evidence of voids or other internal defects.

c. The three sample absorber plates will be subjected to an alpha count to determine if any possible surface contamination exists. The plate shall be checked for alpha contamination by means of gas flow proportional counting or a similar method. Alpha contamination equivalent to 0.5 micrograms of U-235 per square foot shall be the maximum allowable level of contamination. It is assumed that the one microgram of U-235 per square foot is equivalent to 150 disintegrations per minute per square foot of plate surface.

d. Homogeneity - Two of the three sample plates shall be examined for homogeneity. Five miniature samples of full-plate thickness, approximately one square inch in area, shall be extracted from each of the two plates on a diagonal between core corners and approximately equally spaced beginning 2 inches from core-end and 1/4 inch from core-edge interfaces. These samples shall be dissolved and chemically analyzed for total europium content, and the results expressed on a weight per cent basis. For each of the two plates, variation from location to location shall not exceed 5% of the nominal europium content.

e. Bond Integrity - Five transverse samples, equally spaced along the plate length, and three longitudinal samples from each end, equally spaced across the width, shall be removed from one of the three sample plates. After proper preparation and electrolytic etching with 5% chromic acid reagent, the samples shall show no evidence of lack of bond at the clad-frame interface or at the clad-core and frame-core interfaces upon metallographic examination at 100 diameters.

f. Clad-Core-Clad Thickness - The five transverse samples used in "e" shall be measured to determine clad-core-clad thickness. As measured by calibrated eye-piece, the thickness at all points shall show compliance with specified thicknesses.

g. End Conditions - The six longitudinal samples used in "e" shall be examined metallographically. These samples shall show no evidence of the presence of core material in the inactive portions as dimensionally specified.
h. **Fragmentation and Stringering** – One longitudinal sample at least one-half inch long shall be taken from the core of two of the plates. When examined metallographically, none of these samples shall exhibit fragmentation and stringering greater than that illustrated in Figure 1. The europium oxide shall exhibit no evidence of reaction with matrix, clad, or frame material at magnifications up to 1500X.

**Figure 1. Typical Microstructure of Fabricated Absorber Plate Containing 36 w/o Europium Oxide in Stainless Steel. (As polished 250X.)**

5. **Conformance of Fabrication Procedure**

The exact procedure used in fabricating the sample plates and which is finally proposed to be used in fabricating the absorber plates shall be furnished to the Contracting Agency, or its authorized representative at the conclusion of fabrication of qualification plates and at least one week in advance of the request for approval of qualification. Unless otherwise specifically authorized
in writing by the Contracting Agency, or its authorized representative, the procedure used in fabricating the sample plates shall be strictly adhered to in fabricating the final plates and the absorber sections.

F. Qualification of Procedures for Fabrication of Absorber Section

1. Introduction

In addition to the qualification procedures for fabricating absorber plates, the manufacturer is required to qualify the operator and the procedure to be employed in welding the absorber sections.

2. Method of Qualification

Qualification shall be performed by subjecting one assembled and welded solid stainless steel absorber section to the tests outlined in Section E, item 4. Approval of qualification shall be given by the Contracting Agency, or its authorized representative when it is demonstrated to the satisfaction of the Contracting Agency, or its representative that the requirements have been met.

3. Preparation of Qualification Sample

The simulated absorber shall be prepared in conformance with the exact and complete assembly procedure which is proposed to be used in the manufacture of the absorber sections. Solid stainless plates of specified dimensions are to be used in the qualification section. Approval of qualification must be obtained prior to inception of manufacturing of the absorber sections from the Contracting Agency, or its authorized representative. Approval by the Contracting Agency or its authorized representative of the procedure used in establishing qualification or of any modification thereof will not relieve the manufacturer of any responsibility for any phase of the fabrication of the absorber section or for conformity to specification requirements.

4. Test for Qualification

a. Visual Inspection - The welded test section shall be inspected for over-all cross-sectional width and length dimensions and for weld integrity. The test section shall meet all dimensional requirements. The welds shall be in accordance with good commercial practice. The bead shall be continuous and limited to an area .050 to .200 in. measured from the outside edge of the absorber.

The finished test section must pass through a final test box with internal dimensions 2.650 x 2.650 x 26-1/16 in. without binding at
any point. Any binding of the test section in the test box is cause for rejection.

b. Weld Integrity - Four transverse sections, equally spaced along the length of the finished test section shall be removed and examined metallographically for weld penetration. The weld bead shall be essentially flush with the plate surface. The average depth of penetration should be .025 in. A penetration of less than 0.015 in. as measured from the plate surface by a filar micrometer is cause for rejection. The welds shall be essentially free of porosity, inclusions, and tungsten particles.

5. Conformance of Fabrication Procedure

These specifications cover materials and processes required to produce completed elements. The procedural specifications are specific for manufacturing these components at ORNL and other facilities, and represent a detailed guide for fabricators. It is recognized that because of differences in equipment, fabricators may be required to modify some of the detailed procedures to arrive at the same result. Prior approval for any deviation from these specifications must be approved by the Contracting Agency or its authorized representative.

Delivery of an acceptable product, including materials, dimensions, loading, and all other requirements, remains the responsibility of the manufacturer.

The exact procedure used in fabricating the test section and which is to be used in fabricating the final absorber sections shall be furnished to the Contracting Agency, or its authorized representative at the conclusion of fabrication of the qualification section and at least one week in advance of the request for approval of qualification. Unless otherwise specifically authorized in writing by the Contracting Agency, or its authorized representative the procedure used in fabricating the test section shall be strictly adhered to in fabricating the final assemblies.

G. Liaison and Inspection

Free entry shall be given to the Contracting Agency, or its authorized representative to all areas of the manufacturer's plant at any time during the term of the contract for fabricating the absorber sections. The manufacturer shall provide all reasonable assistance, facilities, and cooperation to the Contracting Agency, or its authorized representative for determination of compliance with specifications or procedure requirements or for inspection purposes as may be required.
The Contracting Agency or its authorized representative will maintain liaison with the manufacturer for the duration of the contract for the following purposes: (1) to provide necessary and reasonable technical assistance as may be required, and (2) to inspect for compliance to the specifications and the approved fabrication procedure. The Contracting Agency, or its authorized representative shall have the right at any time during the term of the contract to reject any and all pieces, parts, components, and products which do not meet the requirements of the specifications, or which have not been fabricated in accordance with the approved procedure, or which fail in any way to meet any of the requirements set forth in this document. Such inspection shall not relieve the manufacturer of any responsibility in any phase of absorber section fabrication or furnishing thereof.

H. Certification

Certification shall be furnished to the Contracting Agency or its authorized representative that all materials used in the fabrication and furnishing of the absorber sections are in accordance with the requirements of these specifications.
II MANUFACTURING PROCEDURES

A. Introduction

The essential operations required in processing the absorber sections are: (1) preparation of the europium oxide, (2) weighing and blending of the compact powders for each absorber subcore, (3) pressing, sintering, and coining into a compact of the required dimensions, (4) assembling of the absorber billet constituents, (5) welding and evacuating the billet, (6) cladding by hot roll bonding, (7) descaling of the hot-rolled plate, (8) flatten annealing, (9) marking and shearing of the composite, (10) machining to final length and width dimensions, (11) assembling and welding of the section, (12) attachment of the handle, (13) cleaning of the weld surfaces, and (14) inspection. After the final inspection, the units are degreased and packaged for shipment to the reactor site. These general procedures and the more specific details, which will be described later, represent methods developed and adopted by the Oak Ridge National Laboratory for manufacturing stainless steel europium oxide absorber sections for service in Core II. It is recognized that, because of differences in equipment, other fabricators may be required to modify some of the detailed procedures to arrive at an equivalent finished product.

B. Records

During processing, positive identification of each absorber plate must be maintained along with appropriate data in order to ensure: (1) proper process control, (2) quality control, (3) metallurgical history record, and (4) removal of rejected material. Records of the following items are maintained and copies furnished to the Contracting Agency or its authorized representative with the finished absorber sections: (1) identification of each lot of prefired europium oxide, (2) master log containing materials make-up data and the detailed processing schedule employed in the manufacturing of each plate (to serve as a guide, the forms of the pertinent records are included in the Appendix) and (3) absorber section inspection record of critical dimensions.

C. Europium Oxide Preparation

Europium oxide obtained from various vendors is normally of low density with extremely fine particle sizes. Such material is not readily suitable for a powder metallurgical dispersoid. Prior to incorporating the oxide in a stainless steel matrix the material is conditioned by a high-firing proc-
ess. The various stages of this operation are as follows: (1) inspection of the received oxide, (2) pressing, (3) firing, and (4) crushing.

1. Inspection of received europium oxide

It is recommended for optimum conditioning properties that the received oxide be obtained in a "dead burnt" form from the oxalate with no additional intermediate temperature treatments. Each batch of received oxide is inspected for total europium oxide and thoria content. The material should contain a minimum of 98% europium oxide.

2. Pressing

Prior to high firing, the received powder is pressed into small cylindrical pellets to facilitate powder handling. Approximately 35 g of the material measured by a 10 cc stainless steel beaker is pressed into an individual pellet. The measured amount of oxide is poured into the cavity of a double-acting, 0.8 in. diameter powder metallurgy die, leveled, and pressed into pellet form by applying a pressure of about 4 ton/in.². The resulting pellets are of sufficient strength for handling and have dimensions 0.8 in. diameter by 0.8 in. length.

3. Firing

Approximately 10 pellets weighing between 300 - 350 g comprise the firing batch size. The individual pellets are placed in a tungsten crucible approximately 2 x 2 x 7 in. in size which is constructed by edge welding 1/8 in. sheet stock. The loaded tungsten crucible is inserted in a reactor grade graphite crucible mounted in the center of an induction coil. Amorphous thermatomic carbon powder is packed between the reactor grade graphite crucible and the outer quartz furnace tube to serve as an insulator. A reactor grade graphite block with an attached off-gassing stem is placed over the reactor grade graphite crucible to prevent the thermatomic graphite from contaminating the europium oxide pellets. The remainder of the quartz tube is then filled with amorphous thermatomic carbon powder and finally capped with a gas-tight, brass end-plug. A hydrogen gas stem is attached to the lid to permit passage of hydrogen to the pellets during the firing operation. The exit hydrogen gas escapes through the stem attached to the top of the graphite crucible. The pellet firing is carried out at 3090°F for a period of three hours. The hydrogen atmosphere gas is dried hydrogen gas with a dew-point of at least -60°F. The power is varied initially to control the rate of temperature rise in the furnace; a rate of 570°F/hr is average. Temperature measurements are made on the surface of the pellets by sighting an optical pyrometer through the off-gassing stem. Upon completion of the firing, the power is shut off and the furnace air cooled to approximately 300°F at which time it is disassembled and the fired pellets removed.
Fig. 2. Furnace Assembly for High-Firing Europium Oxide Pellets.
4. **Crushing**

The fired pellets are crushed to -325 mesh size in a mortar and pestle using a series of intermediate screening operations. Initially one-third of the fired batch size is placed in a mortar and crushed using only impact-type strokes. Circular grinding motions were found to produce considerable fines in the crushed oxide. After the pellets have been initially broken down, the product is placed in a series of vibrating screens which separate out the +100, -100 +270, -270 +325, and -325 mesh fractions. Each fraction is then recrushed until the entire material is of a -325 mesh size. The various sizing separations are necessary to minimize presence of fines. The final product from each batch is examined macroscopically to determine if the particles are irregular and not rounded. Since the percentage of europium generally varies in high firing, a representative sample of each batch of hi-fired oxide is analyzed by wet chemistry for total europium content expressed in weight percent.

A specimen from each processed lot or batch of europium oxide shall be tested for weight change by firing in a dry hydrogen (dewpoint at least 
-60°F) atmosphere for 1-1/2 hours at 2250 ± 25°F. The specimen shall be cold pressed in a 1/4 inch diameter die at a pressure of 3100 psi before firing. The pellet is fired on a molybdenum lined type 316 stainless steel boat in a leak tight inconel muffle. Any weight gain or a weight loss in excess of 0.008 gms will be cause for rejection of the lot or batch of europium oxide which this specimen represents. The specimen shall weigh before firing.

After high firing the europium oxide must be shipped, stored and handled under conditions which preclude the pickup of any moisture.

D. **Absorber Core Manufacturing**

1. **Calculations**

The neutron absorber and matrix powders are specified in terms of europium and stainless steel. Since the poison used is in the form of a complex europium oxide, it is necessary to determine by calculation the quantity of this material to be incorporated into each core. Each batch of hi-fired europium oxide is assayed for total europium content. Since elemental powders are employed as the matrix material, the quantity of each constituent must be determined by calculation.

Sample calculations illustrating the method utilized in determining the quantities of europium oxide and matrix powders required in the made-up of each absorber core are listed below:
(a) Data Required

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt % Eu in europium oxide</td>
<td>85 average value</td>
</tr>
<tr>
<td>Grams of Eu per plate</td>
<td>151</td>
</tr>
<tr>
<td>Wt % Fe in matrix steel</td>
<td>71</td>
</tr>
<tr>
<td>Wt % Ni in matrix steel</td>
<td>11</td>
</tr>
<tr>
<td>Wt % Cr in matrix steel</td>
<td>18</td>
</tr>
<tr>
<td>Dimensions of absorber core in finished plate</td>
<td>20.75 x 2.213 x 0.090 in.</td>
</tr>
<tr>
<td>Densification of core in finished plate</td>
<td>0.957</td>
</tr>
</tbody>
</table>

(b) Determination of Grams of europium oxide per plate

\[
\frac{151 \text{ g of Eu} \times 1.00}{85\% \text{ of Eu in europium oxide}} = 177.65 \text{ g}
\]

(c) Determination of Grams of Matrix Stainless Steel per Plate

1. Required core volume in finished plate:

\[
20.75 \times 2.213 \times 0.090 \times 16.38 = 67.69 \text{ cm}^3
\]

2. Volume of core occupied by wrought stainless steel foils*:

\[
2.103 \times 2.275 \times 0.005 \times 4 \times 16.38 = 1.57 \text{ cm}^3
\]

* Values based on initial core dimensions.

3. Core volume occupied by europium oxide bearing dispersion

\[
67.69 \text{ cm}^3 - 1.57 \text{ cm}^3 = 66.12 \text{ cm}^3
\]

4. Volume occupied by dispersion materials initially:

\[
66.12 \text{ cm}^3 \times 0.957 \text{ (densification)} = 63.28 \text{ cm}^3
\]
(5) Material volume occupied by europium oxide based on europium oxide density = 7.4 g/cm$^3$

\[
\frac{177.65 \text{ g}}{7.4 \text{ g/cm}^3} = 24.01 \text{ cm}^3
\]

(6) Material volume occupied by matrix steel:

\[
63.28 \text{ cm}^3 - 24.01 \text{ cm}^3 = 39.27 \text{ cm}^3
\]

(7) Determination of grams of matrix steel per plate based on stainless steel = 7.9 g/cm$^3$

\[
39.27 \text{ cm}^3 \times 7.9 \text{ g/cm}^3 = 310.23 \text{ g}
\]

(8) Determination of grams of matrix steel constituents per plate:

(a) Fe - 71 wt %

\[
310.23 \text{ g} \times 0.71 = 220.26 \text{ g}
\]

(b) Ni - 11 wt %

\[
310.23 \text{ g} \times 0.11 = 34.13 \text{ g}
\]

(c) Cr - 18 wt %

\[
310.23 \times 0.18 = 55.84 \text{ g}
\]

(9) Determination of total grams of core constituents per plate:

\[
177.65 + 220.26 + 34.13 + 55.84 = 487.88 \text{ g}
\]

(10) Determination of weight percent europium oxide per plate:

\[
\frac{177.65}{487.88} \times 100 = 36.41\%
\]

(11) Since each absorber core was manufactured in three parts, the grams of europium oxide per sub-core are:

\[
177.65 \text{ g} \times \frac{1}{3} = 59.127 \text{ g}
\]
Likewise the grams of Fe, Ni, and Cr in the matrix steel are:

\[
\begin{align*}
\text{Fe} & \quad 220.26 \times \frac{1}{3} = 73.42 \text{ g} \\
\text{Ni} & \quad 34.13 \times \frac{1}{3} = 11.38 \text{ g} \\
\text{Cr} & \quad 55.84 \times \frac{1}{3} = 18.61 \text{ g}
\end{align*}
\]

As might be expected, the densification factor \( \frac{\text{measured volume}}{\text{theoretical volume}} \) of the fabricated core material has a significant effect on the amount of elemental stainless steel powder required to meet dimensional specifications in the finished composite plate. This factor has been determined to be approximately 0.96 for the specific equipment and processing methods employed at the Oak Ridge National Laboratory. However, any major change in equipment or procedure will probably shift this factor; and under such circumstances, it will be necessary to alter the grams of elemental stainless steel required in the material charge for the absorber compact. Such changes in the total material charge affect the grams of elemental stainless steel powder in the three sub-cores which are stacked in the billet assembly to make up the total core material charge.

2. **Weighing of Component Powders**

The absorber section consists of three compacts or sub-cores stacked on top of each other. The component powders for each sub-core are separately weighed and then combined in a single blending jar. With the exception of possible losses during subsequent pressing and sintering operations, this method offers accurate accounting and reproducibility of the critical ingredient, europium oxide, in each absorber sub-core and ultimately in each absorber core within the limits of the accuracy of the weighing balance. Europium oxide is the first material loaded into the blending jar and is followed by the constituents of the elemental stainless steel powder, namely electrolytic iron, nickel, and chromium. Since the absorber section contains no contaminated or active materials, handling of all powders can be done in a conventional manner. During processing, five jars comprise a batch. The individual weighing procedures are as follows:

(a) **Weighing of europium oxide** -

The neutron poison, europium oxide, in the sub-core is weighed to an accuracy of at least 0.03% on a Grammatic balance of 200 g capacity. A 4 x 4 in sheet of glazed paper with glazed side up and of known weight is placed on the pan. The europium oxide is added to the paper and accurately weighed. The material is then poured into a clean, dry, wide-mouth, glass jar of 4-ounce capaci-
ity. A camel's hair brush is used to brush any remaining particles of europium oxide into the jar, which is then capped.

(b) Weighing of the elemental stainless steel powder -

The elemental stainless steel constituents, electrolytic iron, nickel, and chromium are weighed in exactly the same manner as the europium oxide, with the exception that a triple-beam balance is used. The individual powders are weighed to a tolerance of ± 0.01%. After each weighing, the individual powder is then transferred to a blending jar containing the previously weighed europium oxide. The jar is immediately recapped. This procedure is again repeated for the two remaining constituents.

These operations are repeated until the lot of five jars has been processed. After final weighing and capping, the joint between the cap and the jar is sealed with masking tape, and each jar identified by europium oxide batch number.

3. Blending

The powders are blended to achieve a homogeneous mixture of europium oxide and elemental powders. A modified U.S. Stoneware Company Double Cone Blender, Model 733, is used. The blender cones are replaced by a pair of two-quart steel cans mounted on the motor shaft at an angle of 30 degrees with the vertical. Ten jars or two lots, each containing the specified quantities of europium oxide and elemental powders, are loaded into each can with sufficient padding to prevent breakage of the jars during the blending operation. The cans are rotated at this oblique angle for three hours.

4. Initial Cold Pressing of the Core Ingredients

The initial operation in shaping the blended powders into a compact suitable for assembling into an absorber plate billet is to cold press into a "green" compact. A Baldwin press of 150-ton capacity and with a Vickers Hydraulic control system to permit variable movement of the ram is employed for compaction. The press has two rams, an upper fixed ram extending down from the top crosspiece and a lower ram which is movable. On the top of the lower movable ram is fastened a large platen onto which is placed the powder metallurgy die set. The die set consists of a female die which is spring mounted to the movable ram and a die punch and filler block. The bottom of the die punch contacts the movable ram, while the top of the filler block, inserted into the die cavity after the powder, contacts the stationary ram.

After assembly has been completed, a solution of 10 wt. % C. P. Stearic Acid - 90 wt. % carbon tetrachloride, which serves as a die lubricant for each
die charge, is applied with a 1/4 inch camel's hair brush around the top of
the die cavity and the lower sides of the filler block. The die face is oc-
casionally lubricated when the pressed core adheres to it because of wear.
The blended powder from each jar is poured into the die cavity. The jar is
brushed thoroughly to ensure complete transfer of all powders. Once the
core material has been loaded into the cavity, it is leveled with the straight
edge of a scoopula. The filler block is inserted into the die and the lower
ram then raised until the die insert contacts the stationary ram. The blended
powders are initially pressed at 10 tsi pressure (approximately 45-tons total
load) for 15 seconds to a thickness of approximately 0.404 in. The filler
block is removed and the "green" compact ejected by elevating the die punch
with the hand lever. Extreme care must be exercised in removing the "green"
compact to avoid breakage since the compact is of low strength and density.
The pressed compact is then carefully placed on the sintering boat to mini-
mize handling.

5. Initial Sintering of "Green" Compacts

The sintering operation is carried out in a General Electric 20-kw labo-
atory molybdenum-wound furnace equipped with a 3 in. diameter Inconel muf-
ble. Thirty-six inches of the muffle extends beyond the furnace and acts as
a cooling chamber. The sintering temperature is 2250°F. Uniformity of
temperature along the length of the sintering boat is maintained at ± 25°F.
Dry hydrogen with a dewpoint of -60°F, as measured by an Accurate Dewpoint-
er at the gas inlet, is used as atmosphere in the muffle. Hydrogen flow required
for this size muffle is approximately 20 ccf under steady-state conditions,
although the gas flow is increased when the muffle door is opened during inser-
tion or removal of compacts. The sintered compacts do not exhibit any evidence
of oxide formation.

The sintering boat which contains the pressed "green" compacts is con-
structed of type 316 stainless steel formed into the shape of an "H". The
crossbar between the two vertical sides is 2-1/2 in. wide x 14 in. long, and
is located approximately 1/2 in. above the bottom of the legs.

Prior to positioning five "green" compacts end to end along the length of
the boat, a molybdenum-foil liner is inserted to prevent the sub-cores from
contacting the stainless steel. This liner has 1/16 in. diameter molybdenum
wire spacers welded along its length upon which the five compacts rest. This
spacer wire is used to permit access of hydrogen gas to all surfaces of the
compacts.

The sintering boat containing the five compacts is inserted in the furnace
at 2250°F and the system purged with hydrogen. The compacts remain in the
hot zone for 1-1/4 hours. The boat is moved to the cooling chamber, allowed
to cool to 450°F under the hydrogen atmosphere, removed from the muffle,
and air cooled to room temperature. Again, in an effort to minimize handling, cores are not removed from the boat until re-inserted into the pressing die for additional compaction.

6. Repressing the Sintered Compacts

The sintered compacts are repressed in the original die set described in Item 4, Section D. The shrinkage encountered during sintering permits re-insertion of the compact into the same die cavity. After careful insertion, the sintered compacts are pressed under a pressure of 31 tsi (approximately 150-ton load) for 15 seconds to improve densification. The repressed core thickness is approximately 0.339 in. The repressed compacts are ejected from the die, and repositioned on the sintering boat.

7. Resintering of Compacts

The repressed compacts are resintered in the furnace described in Item 5 of Section D under dry hydrogen for 1-1/4 hours at 2250°F and cooled in cold leg of muffle to 570°F. This second sintering promotes densification and alloying of the elemental powders.

8. Coining of Sintered Compacts

The shrinkage encountered during the second sintering operation permits re-insertion of the compact into the original die cavity for coining. After careful loading, the resintered compacts are pressed under a pressure of 31 tsi for 15 seconds to obtain the desired dimensions and improved densification. The coined compacts are ejected from the die and transferred to the inspection area. The coined compact thickness is approximately 0.327 in.

9. In-Process Inspection

Since it is necessary to control the europium content of the absorber, it is desirable to weigh the compacts prior to the completion of the sub-core processing. Because of uncertain losses due to the reduction of metal oxides by hydrogen during sintering, it is impractical to consider weight losses of the compact after this operation. The handling losses are determined therefore prior to loading the furnace for initial sintering. For convenience, all five sub-cores are weighed together. If the total permissible deviation for a single core is exceeded by the entire batch, each sub-core must be individually weighed for acceptance or rejection. Deviation from the charged weight greater than specified in Section V-A for each core is the basis for rejection. Compacts with obvious chips and flaws, of course, are weighed individually and not included in the batch weighing. After sintering the compacts are examined to determine whether any obvious chipping or spallation occurred during the required handling.
Each core is measured with a micrometer after processing to determine if the compact meets the thickness requirement of 0.327 ± 0.002 in. Length and width dimensions are established by die design.

10. **Core Storage**

Each acceptable lot of cores is wrapped in paper. These packages are marked to identify batch number and europium oxide lot number, and inserted in an ordinary desiccator if storage time is to be less than 48 hours. If a storage time of more than 48 hours is required, a vacuum desiccator is used.

E. **Billet Assembly**

1. **Material Preparation**

The billet containing the europium oxide stainless steel dispersion is designed to permit evacuation of the billet interior prior to rolling into a composite plate. The components consist of two wrought stainless steel cover plates; a frame into which the europium oxide compacts are inserted; three sub-cores containing the europium oxide; and five stainless steel foils.

   a. **Stainless Steel Cover Plates** – The cover plates are fabricated from 5/16 in. plate (minimum thickness of 0.318 in.) which may require cold rolling to the specified thickness of 0.320 in ± 0.002 in. The plates are sheared to 4-3/4 in x 4-3/4 in ± 1/16 in, which allows approximately a 3/16 in. overlap of the cover plate on the frame. Since only very small reductions are necessary during cold rolling, it is recommended practice to shear the received plate to the above dimensions and to reduce to the required thickness by cross-rolling.

   b. **Picture Frames** – Picture frames are machined from 1-in. plate. Since the billet is to be subsequently evacuated prior to rolling, holes are drilled into the frame to provide a free path for evacuation. It is important that the finished plate be free of potential defects caused by the drilled holes. By the design, in shearing and machining the plate to final size, these regions will be in the scrapped portion of the plate. An entrance hole, 0.225 in. in diameter, is drilled in the geometric center of the tail edge of the plate frame to a depth of 3/8 in. A corresponding 0.225 in. diameter hole is drilled through the frame perpendicular to and meeting the end of the first hole to form a "T".
c. Evacuation Tube — The tube for evacuation of the billet interior is type 304 or 347 stainless steel, with a nominal outside diameter of 0.225 in., an inside diameter of 0.175 in. and a length of 12 in.

d. Foils — Foils are prepared from electrolytic iron, chromium, and nickel powders, in the weight ratio of 71 - 18 - 11, respectively. The silicon content of each powder is limited to a maximum of 0.03 wt. %. The foils serve two purposes: (1) as a bonding aid between the three sub-cores and (2) as a barrier to prevent possible reaction between the europium oxide and the wrought stainless steel. The foils are prepared by blending the specified quantity of each of the elemental powders for 3 hours in an oblique blender; cold pressing at 31 ksi; sintering in dry hydrogen of -60°F dewpoint; at 2250°F for one hour; and coining at 31 ksi. The coined stainless steel is cold rolled on a Bliss 4-high mill to 0.005 in. ± 0.0005 in. with intermediate one hour anneals at 2250°F to minimize edge cracking. Light reductions per pass coupled with cross-rolling yield the most satisfactory material.

2. Assembly and Welding of Billets

Immediately prior to billet assembly, both surfaces of the picture frame and the surfaces of the cover plates which contact the picture frame and core are thoroughly vapor degreased and surface oxide is removed by scratch brushing with a power-driven stainless steel brush. The evacuating stem is inserted approximately 1/8 in. into the hole provided in the frame and welded to the frame with type 347 stainless steel filler rod. The tube is purged with an inert gas during welding to prevent melting of the tube wall.

The billet is assembled by stacking the three sub-compacts in the frame hole. Between each sub-compact a foil is placed to improve the compact to compact bonding during hot rolling. To prevent contact of the compacts with the frame and potential reaction of europium oxide with wrought stainless steel, foils are placed around the periphery of the frame hole prior to insertion of the stack of sub-compacts. Foils are placed at the cover plate-compact interfaces and held in place by bending the edges of the foil over the sub-compact. After insertion of the compacts, cover plates are placed on top and bottom of the frame and held in place by "C" clamps. The cover plates overhang the frame by approximately 3/16 in. on all sides to facilitate welding.

Cover plates are attached to the frame by heliarc welding. Two heavy weld passes employing a type 347 stainless steel filler rod are made completely around the periphery of the two frame-cover plate joints. All welds are brushed and examined for possible flaws. The sealed billet is subsequently leak tested under water with a 20 pound internal helium pressure applied through the evacuation stem. The billets are then evacuated to a final vacuum.
of less than 10 μm as indicated on a thermocouple vacuum gauge. During this operation, the billet is periodically heated to approximately 500°C to drive off any water vapor and entrapped gases. The intermittent heating is continued until there is no further indication of loss in vacuum upon re-heating at which time the evacuation stem is sealed by hot forging.

F. Composite Plate Fabrication

1. Hot Rolling

The assembled and evacuated billets are hot rolled in lots of 2 to a thickness of 0.156 in. ± 0.002 in. on a two-high Mesta mill equipped with 20-inch diameter x 30 in. wide rolls. However, the lot size is determined by the muffle available for heating the billets. A furnace capable of maintaining 2100°F ± 25°F over a muffle length of 40 in. is recommended. The muffle is purged from the rear with hydrogen gas with an inlet dewpoint of less than -50°F as measured by an Accurate Dewpointer. A flow rate of 250 cfh is generally used.

The billets are heated to 2100°F for at least 90 minutes prior to the first pass, and are reheated for at least 5 minutes between subsequent passes. The reheat time is reduced to 3 minutes between each pass near completion of the hot rolling. A second lot of billets is introduced into the furnace for preheating at the initiation of rolling the preceding billets to allow continuous hot rolling. After the final pass, the plates are replaced in the muffle for a 5 minute anneal, and are then air cooled.

On all billets, the side containing the hot forged evacuation stem is gripped so that the opposite end can enter into the rolls. This practice is continued throughout the hot rolling to assure a single rolling direction, although the billet is rotated 180 degrees about its longitudinal axis between passes. The hot rolling is performed in approximately 22 passes using 10% reductions in thickness during each pass as indicated by mill settings. Lighter reductions are made, as required, to roll to 0.156 in. ± 0.002 in. near end of fabrication schedule. The plate thickness is measured over the core area with a micrometer to determine the actual thickness of the hot-rolled plate. As the thickness of the plate approaches 0.170 in., core length is measured between subsequent passes with a tape to ensure that the plates are not rolled over-size in length. The core length can be readily measured in this manner since the dispersion-type core can be delineated from the wrought stainless steel by a difference in heat color.

Special care is exercised during rolling to minimize cambering or "rainbowing" of the plates which tends to decrease the amount of stainless steel along the inactive edge in the final machined plate. Cambering of the
absorber plate may possibly be corrected, depending on severity, by inserting the plate through the mill at a slight angle. A level mill and proper feeding are recommended to circumvent this difficulty.

After cooling and prior to end trimming the hot-rolled plates are numbered consecutively starting with "1" for the first plate processed. Numbers, one-half inch high, are stamped on the inactive section at the end opposite the evacuation stem or the leading end of the plate. Care must be exercised to insure that this identification is maintained during subsequent processing operations. Each number is entered into the record.

2. Pickling of Hot-Rolled Plates

After the plates have been properly identified, they are sheared on a power-driven shear to a total length of 34 inches. The trailing end of the plate is sheared to within 1-1/2 in. of the core and the remaining wrought stock removed from the leading plate end. The oxide scale which formed during hot rolling and cooling is removed by pickling the plates in an aqueous solution of 5% HF and 15% HNO₃. After pickling, the plates are thoroughly washed with water to remove all traces of acid. The plates are then dried and visually examined for surface pitting, oxide scale inclusions, and other possible defects.

3. Flatten Annealing of Absorber Plates

To prevent warpage of the absorber plates during subsequent machining and shearing operations, it is necessary that the composite plates be flat and fully annealed. The use of flat plates also facilitates assembling into the required box array. Both the flattening and annealing of the absorber plates is accomplished in a single operation.

Each cleaned plate is covered on one side with a thin coat of a mixture containing one part by volume of Fisher 5F Precisionite levigated alumina and ten parts of water. A three-inch camel's hair brush is used to apply an even coating. The coated plates are allowed to dry for at least 15 minutes. The jig is composed of two platens for clamping the absorber plates together. Six absorber plates are stacked together with the coated side adjacent to the uncoated side. The stack is placed between the platens, and the clamping bolts are firmly tightened. The loaded assembly is dried in an oven at 330°F for a minimum of 16 hours.

Annealing is accomplished within a leaktight Inconel muffle inserted into a 56-kw Globar furnace at a temperature of 1832°F ± 25°F. The muffle has a cross-sectional dimension of 7 x 9 in. and is 6 feet in length. Bright annealing of the plates is obtained under a purge of dry hydrogen with a dew-point of -80°F as measured by an Accurate Dewpoint Measuring Instrument at the furnace inlet.
Furnace temperature at insertion of the loaded platen assembly ordinarily does not exceed 570°F. The muffle is purged with dry helium prior to insertion of the plates and during the period between 570°F and 1200°F. At 1200°F the helium atmosphere is replaced by dry hydrogen. A flow rate sufficient to bright anneal stainless steel is recommended; 240 cfh is generally used. The temperature is increased to 1832°F at a rate of approximately 540°F per hour. The plates are held at 1832°F for two hours, and then slowly furnace cooled to 570°F. At this temperature, the hydrogen atmosphere is replaced with a dry helium purge; the assembly removed from the furnace; and air cooled. After disassembly, the absorber plates are scrubbed under flowing water to remove the coating of alumina, air dried and inspected again for defects.

4. Preliminary Machining of Absorber Plate

To assure a true plate reference edge for radiographically locating the absorber core, it is necessary to machine one edge of the plate. A stack of six plates is edge aligned on the bed of a milling machine and held in position by bar clamps. A series of skim cuts are made to a total depth of approximately 1/4 in. or until smooth, uniform plate edges are obtained. The edges of the plates are filed to remove machining burrs and the plates are cleaned by vapor degreasing.

5. Radiography

The edge machined plates are radiographed to delineate the core in the composite plate for subsequent marking. The plates are positioned on cassettes containing a fine-grained film such as Eastman type "M" and exposed to an X-ray source positioned 54 inches above the cassette for four minutes. Without using lead filters and with an X-ray source power setting of 130 KV and 10 m.a., suitable radiographs are obtained. Normally two radiographs of the plate are taken; the first to obtain over-all dimensions and the second to obtain a clear radiograph of the trailing end of the core.

6. Marking of Plate for Shearing and Final Machining

Measurements at three locations along the core length are made and transposed to the plate delineate the core on the plate surface. The measurements are taken with reference to the machined edge of the plate. After the position of the core is located, the resulting core width is subtracted from the required plate width and the difference equally divided and marked at the three reference locations. These markings serve as the width boundaries during machining to final size. The location of the trailing end of the core is critical since the engineering drawing specifies a minimum of 1/4 in. of inactive stainless steel at this end. The end is located from the radiograph and a mark scribed 1/4 in. from the core end.
this scribe mark, the final plate length of 26-1/16 in. is measured to the opposite end of the plate and a mark scribed.

After the machining reference lines are scribed, additional lines 1/8 in. outside of the machining lines are located at the ends and along both sides. These lines represent the sheared length and width dimensions. The identifying number is again stamped at the leading end of the plate which contains the nominal 5 in. length of inactive stainless steel within the machining line boundaries.

7. Shearing of Plates Prior to Machining

To minimize the amount of wrought stock removal prior to machining, the plates are sheared to within 1/8 in. of the final machining dimensions. The plates are sheared on a power-driven shear suitable for shearing 3/16 in. thick stainless steel. In shearing to width dimensions, the plates are positioned and clamped to the shear table with three bar-type clamps equally spaced along the plate length. Clamping is not required in shearing the plate ends.

8. Inspection for Core Alignment

Prior to machining the absorber plate, it is recommended that the core alignment be inspected. The 1/8 in. excess wrought stock provides ample room for re-alignment if required. The core alignment is inspected by radiographic examination. Radiographs are obtained in the same manner as described in Section F, Item 5, except that four plates are positioned on the cassette at one time.

9. Machining of Absorber Plates to Final Dimensions

In this operation, it is recommended that a single plate be machined at one time. Each plate is first machined to the final width dimensions. The sheared absorber plate is placed on the table of a vertical milling machine. A 3/8-in. aluminum plate is positioned between the absorber plate and the table to elevate the absorber plate. Using a dial gauge indicator, the machining scribe marks are aligned with the traverse of the table and held in position by bar-type clamps at the plate ends. Three additional bar clamps are positioned and firmly locked along the plate length in such a manner as to permit machining along one side of the plate. After checking the plate alignment, the plate edge is machined to the scribed line. The three bar clamps are then removed without disturbing the plate and re-positioned on the opposite side of the plate. After rechecking the plate alignment, the remaining plate edge is machined to the scribed line. The width dimension is inspected with a micrometer.
The end of the plate at which the minimum 1/4-in. inactive stainless steel end is specified is likewise individually machined to the scribed machining mark. This is accomplished after machining the second plate edge by leaving the plate locked in position and removing the appropriate end bar clamp. Six plates which have been machined at the critical end are stacked together, butted against an end plate and gang milled to final length.

After completion of the machining operation, the plates are degreased in a vapor degreaser and the machined edges are lightly deburred by hand filling at a 45 degree angle.

10. **Final Inspection of Plates**

a. All absorber plates are inspected to ensure that they meet final dimensions. The over-all length, width, and thickness measurements are made with a calibrated straight edge and micrometers, respectively. The finished machined plates are then radiographed to delineate the inactive stainless steel. Measurements are made from the radiographs to determine acceptance or rejection. Plates with less than 0.080 in. inactive stainless steel at any point along the edges are rejected. Plates with less than 3/16 in. at the end specifying a nominal 1/4 in. inactive stainless steel are also rejected.

b. Contamination Check. The absorber plates will be subjected to an alpha count to determine if any possible surface contamination exists. The plate shall be checked for alpha contamination by means of gas flow proportional counting or a similar method. Alpha contamination equivalent to 0.5 micrograms of U-235 per square foot shall be the maximum allowable level of contamination. It is assumed that one microgram of U-235 per square foot is equivalent to 150 disintegrations per minute per square foot of plate surface.

G. **Manufacturing of the Absorber Section**

Immediately prior to assembly, all components are visually inspected. The absorber plates are again degreased to ensure cleanliness.

1. **Assembly Jigs**

Two jigs are required for the assembly of the absorber section. The initial assembly is performed around a solid graphite block. A 1/2 in., 45 degree cut is machined from all corners of this block to prevent contact of graphite and stainless steel during subsequent welding. Final assembly of the section is achieved using a split graphite block. This split block wedge design allows convenient removal after final welding.
2. Assembly and Welding of the Section

The following sequence is recommended for the assembly and welding of the absorber section.

(a) Plate No. 1 serves as the base plate and is positioned on two 1 x 2-in. parallel bars equally spaced on a surface plate. The plate end opposite the identifying number is butted against an angle plate.

(b) The solid graphite block is positioned on plate No. 1 and butted against the angle plate. The right side of the block should lie approximately 0.156 in. inside the edge of the base plate.

(c) Plate No. 2 is placed perpendicular to the surface plate with the edge butting the parallel bars and the inner surface butting the edge of plate No. 1. Care is taken to assure that the identifying number is opposite the end of the plate adjacent to the angle plate.

(d) Plate No. 3 is placed perpendicular to the surface plate with the edge butting the inside of plate No. 1. To hold the plates in position, "C" clamps are spaced along the length of the assembly. Again, care is exercised to assure that the end opposite the identifying plate number is adjacent to the angle plate.

(e) Plate No. 4 is then placed on top of the graphite block. The right edge of the plate is butted against the inner surface of plate No. 3 and the inner surface butting the edge of plate No. 2. The position of the numbered end of the plate is again checked to assure proper positioning adjacent to the angle plate.

(f) After inspecting the position of the identifying numbers on each plate, the assembly is rigidly clamped together with 5-in. "C" clamps. The end of the section with the 1/4-in. nominal inactive stainless steel is squared against the angle plate. The width and height of the assembly is measured with a micrometer. If dimensions are less than nominal, shims are placed at the proper points to increase the dimensions to 2.619 in. ± 0.010 in.

(g) After proper dimensions have been established, the joints between plates No. 1 and No. 2 and No. 1 and No. 3 are heliarc welded. These joints are first tack welded at the corners. The welds are made in conformance with the procedure previously established in qualification. Three-inch-welds are initiated at the corners and continued inward and are staggered from diagonally opposite corners until the entire joints are completed. No filler rod is used during the welding operation.
(h) Upon cooling to room temperature, the "C" clamps are removed from the assembly and the top plate (No. 4) and the solid graphite jig are removed.

(i) The split, wedge-shaped jig is inserted into the partially welded absorber section. The two wedge-shaped blocks are fitted together to form a rectangle with over-all dimensions of 1.80 x 2.307 x 30 in. Since the width of this jig is 1/2 in. narrower than the solid graphite jig, it is centered in the partially welded section with a 1/4-in. clearance between the sides of the jig and the absorber plates.

(j) The top plate or plate No. 4 is replaced on top of the split jig with its right edge butting the inner surface of plate No. 3 and its inner surface butting the edge of plate No. 2. Care is taken to identify the position of the numbered plate end. After squaring the end of the section with the 1/4-in. erective edges, it is secured in place with 5-in. "C" clamps. The dimensions of the assembly are carefully checked with a micrometer to insure conformance to drawing.

(k) The remaining plate joints between plates No. 2 and No. 4 and No. 3 and No. 4 are then heliarc welded in the same manner as described in Item G.

(l) While the welded absorber section is still warm, the clamps are released and the split-graphite block is removed by gently tapping an end of one of the wedges.

(m) All welds are brushed and inspected for weld defects.

3. Attachment of Handle

Prior to attachment of the handle, the entire inner surface of the absorber section is scratch brushed with a long-handled stainless steel brush to remove graphite particles and surface oxide. A hole is drilled through one side of the stainless steel absorber section in accordance with the absorber assembly drawing. The end containing the identifying plate numbers is centered on a drill press table and a 3/8-in. diameter hole drilled through the two absorber plates. A holder pin machined to specifications is then placed in the drilled hole, with approximately a 1/16-in. recess on both absorber plate outer surfaces. The pin is attached by heliarc welding, using type 347 stainless steel filler rod.
H. Cleaning of Absorber Section

The assembled absorber section is cleaned to remove all excess weld metal and discoloration. Removal of the excess weld material and discoloration is accomplished by abrading the entire outer surfaces of the section with fine-grained emery paper or a rotary stainless steel brush. The edges of the absorber are then rounded to a radius of approximately 1/32 in. by hand filing. Upon completion of the mechanical cleaning, the absorber is wiped with a cloth containing a small amount of kerosene to remove the fine metal filings. The kerosene is removed by degreasing in a vapor degreaser.

I. Numbering Absorber Sections

A number 5/16 in. in height is stamped on each plate 1 in. from the end of the absorber which has the nominal 5-in. length of inactive stainless steel. The absorbers are numbered serially, starting with No. 1 and progressing to sequentially higher numbers. A wooden block insert, positioned in the end of the absorber to be marked, prevents possible warpage or deforming of the section during stamping. The appropriate records are entered into the record.

J. Visual and Dimensional Inspection

The assembled absorber is inspected visually for possible flaws such as pits, surface scratches, indentations, and improper welds in accordance with qualification requirements.

The cross-sectional width, length and squareness of each absorber is dimensionally inspected. The cross-sectional width of the absorber is measured with a 3-in. micrometer at five locations along the length. Two measurements, 90 degrees apart, are taken along the center line of the section at distances of 1, 6, 13, 19 and 26 in. from the critical end of the absorber. The measurements of each absorber are recorded on an appropriate inspection record. Absorbers with width dimensions outside the limits of 2.619 in. ± 0.010 in. are rejected.

The absorber length, measured with a calibrated straight edge, must be within the required specification of 26-1/16 in. ± 1/16-in.

The absorber is checked for squareness on a surface plate. Any evidence of twist or other manifestation of out-of-squareness requires that the section be subjected to insertion through the final inspection box. This box is designed with internal dimensions of 2.650 x 2.650 in. and 26-1/16 in. long. Binding at any point during this test is cause for rejection.
K. Preparation for Shipment

Prior to storage and/or shipment, the absorbers are vapor degreased and enclosed in a plastic sheath which is thermally sealed to protect the component. The absorbers are packaged in shipping containers which are designed to prevent damage during shipment.
III APPENDIX

The following forms are illustrated to serve as a guide in helping potential fabricators maintain proper metallurgical history, and other pertinent data requiring permanent record:

1. Rare Earth Powder Metallurgy Core Order Form
2. Powder Metallurgy Fabrication Form
3. Absorber Plate Fabrication Record
4. Radiographic Inspection Sheet
5. Absorber Assembly and Inspection Sheet
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RARE-EARTH POWDER METALLURGY CORE ORDER FORM

Core Order # ___________________________ Order Date __________

No. of Cores Ordered ___________________________ Order Completed __________

Rare-Earth Specifications per core:

1. Type material ___________________________
2. Identification ___________________________
3. Weight per cent _________________________
4. Grams per core __________________________
5. Particle size ____________________________

Matrix Material Specifications per core:

1. Type material ___________________________
2. Base constituents _________________________
3. Identification of Constituents _____________
4. Weight per cent _________________________
5. Grams per core __________________________
6. Particle size ____________________________

Theoretical Total Core Weight, grams ________________________________

Desired Core size ________________________________

Special Instructions _____________________________________________
________________________________________________________________
________________________________________________________________
________________________________________________________________

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RARE-EARTH POWDER METALLURGY CORE ORDER FORM

Data for Core Order # ____________________________

A. Blending Operation

1. Identification of blending container: ____________________________

2. Blending device to be used: ____________________________

3. Blending time desired: ____________________________

Special Instructions: ____________________________

B. Core Processing Operations

1. Initial cold pressing operation: ____________________________
   a. Pressure ______________ tsi
   b. Load ______________ T

2. Initial sintering operation:
   a. Atmosphere ___________, dew point ______________ °F
   b. Sintering time __________ hr, Temperature __________ °F

3. Final pressing operation:
   a. Pressure ______________ tsi
   b. Load ______________ T

4. Final sintering operation:
   a. Atmosphere ___________, dew point ______________ °F
   b. Sintering time __________ hr, Temperature __________ °F

5. Coining operation:
   a. Pressure ______________ tsi
   b. Load ______________ T

6. Other operations: ____________________________
RARE-EARTH POWDER METALLURGY CORE ORDER FORM

Data for Core Order #

Deviations from the specified core composition or fabrication procedure:

Record of individual core weight and thickness:

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<th>Thickness in inches</th>
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Average core Weight ____________________ Average core Thickness ____________________

No deviations from the specified core composition or fabrication procedure are to be made without the prior approval of ____________________
**POWDER-MET FABRICATION FORM**

Batch Number    Lot Number    Type Material    Total Wt Material    Type Cores

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<tr>
<th>No. Cores in Batch</th>
<th>Total Oxide in Batch</th>
<th>Total Batch Wt (A)</th>
<th>Batch Wt After Pressing (B)</th>
<th>Average Wt Loss per Sub-core</th>
<th>Number Cores Accepted</th>
<th>Average Wt Loss per Acceptable Core</th>
<th>Number Cores Rejected</th>
<th>Reason for Rejection</th>
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### ABSORBER PLATE FABRICATION RECORD

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<th>Hot Rolling</th>
<th>Description of Rejection</th>
<th>Miscellaneous</th>
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<td>Date Received</td>
<td>Quality Control Check</td>
<td>Quality O.K.</td>
<td>Minimum Length of Stainless at Critical End (in.)</td>
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ABSORBER SECTION ASSEMBLY AND INSPECTION SHEET

INSPECTION DATA

Cross-Sectional Width Measurements

<table>
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<tr>
<th>Location</th>
<th>Top</th>
<th>Middle</th>
<th>Bottom</th>
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<tbody>
<tr>
<td>Width (A) (In.)</td>
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<tr>
<td>Width (B) (In.)</td>
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Average Width (A) ___ in. + _____ Average Width (B) ___ in. + _____

ASSEMBLY DATA

<table>
<thead>
<tr>
<th>Plate No.</th>
<th>Required Dimensions</th>
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<tr>
<td>1. _______</td>
<td>Cross-Sectional width</td>
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<td>2. _______</td>
<td>Width (A) _______ in.</td>
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<tr>
<td>3. _______</td>
<td>Width (B) _______ in.</td>
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<tr>
<td>4. _______</td>
<td>Length _______ in.</td>
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TEST BOX INSPECTION

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<thead>
<tr>
<th>Acceptable</th>
<th>Slight Bind</th>
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<tr>
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Visual Inspection

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<th>Position</th>
<th>Length Inches</th>
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<td>Edge</td>
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<td>Weld Cracking</td>
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Miscellaneous

ABSORBER FABRICATION RECORD

Type Unit ___________________ Destination ___________________ Shipped _______ Identification Number __________
THIS PAGE WAS INTENTIONALLY LEFT BLANK
## IV DRAWINGS

Absorber Section:

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PART NO.
A9-13-2019

NOTE:
\[\frac{\sqrt{2}}{\sqrt{2}}\text{ ALL OVER.}\]

BREAK SHARP EDGES

FINISH AS INDICATED IN MICRONCHES.

MACHINE FINISH - ROUGH

FLAME CUT OR SAW

ALCO PRODUCTS, INC.
ATOMIC ENERGY DEPT.
SCHENECTADY, N.Y., U.S.A.

PART NO. A9-13-2019

NOTE:
\[\frac{2\sqrt{2}}{2}\text{ ALL OVER.}\]