Standard Specification for Solder Metal

This standard is issued under the fixed designation B32; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ε) indicates an editorial change since the last revision or reapproval.

This standard has been approved for use by agencies of the Department of Defense.

1. Scope

1.1 This specification covers solder metal alloys (commonly known as soft solders) used in non-electronic applications, including but not limited to, tin-lead, tin-antimony, tin-antimony-copper-silver, tin-antimony-copper-silver-nickel, tin-silver, tin-copper-silver, and lead-tin-silver, used for the purpose of joining together two or more metals at temperatures below their melting points. Electronic grade solder alloys and fluxed and non-fluxed solid solders for electronic soldering applications are not covered by this specification as they are under the auspices of IPC – Association Connecting Electronic Industries.

1.1.1 These solders include those alloys having a liquidus temperature not exceeding 800°F (430°C).

1.1.2 This specification includes solders in the form of solid bars, ingots, powder and special forms, and in the form of solid and flux-core ribbon, wire, and solder paste.

1.2 The values stated in inch-pound units are to be regarded as standard. The values given in parentheses are mathematical conversions to SI units that are provided for information only and are not considered standard.

1.3 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to become familiar with all hazards including those identified in the appropriate Material Safety Data Sheet (MSDS) for this product/material as provided by the manufacturer, to establish appropriate safety and health practices, and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:

D269 Test Method for Insoluble Matter in Rosin and Rosin Derivatives
D464 Test Methods for Saponification Number of Naval Store Products Including Tall Oil and Other Related Products
D465 Test Methods for Acid Number of Naval Stores Products Including Tall Oil and Other Related Products
D509 Test Methods of Sampling and Grading Rosin
E28 Test Methods for Softening Point of Resins Derived from Naval Stores by Ring-and-Ball Apparatus
E29 Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications
E46 Test Methods for Chemical Analysis of Lead- and Tin-Base Solder (Withdrawn 1994)
E51 Method for Spectrographic Analysis of Tin Alloys by the Powder Technique (Withdrawn 1983)
E55 Practice for Sampling Wrought Nonferrous Metals and Alloys for Determination of Chemical Composition
E87 Methods for Chemical Analysis of Lead, Tin, Antimony and Their Alloys (Photometric Method) (Withdrawn 1983)
E88 Practice for Sampling Nonferrous Metals and Alloys in Cast Form for Determination of Chemical Composition

2.2 Federal Standard: 4
Fed. Std. No. 123 Marking for Shipment (Civil Agencies)

2.3 Military Standard: 5
MIL-STD-129 Marking for Shipment and Storage

3. Terminology

3.1 Definitions:
3.1.1 producer, n—the primary manufacturer of the material.

3.2 Definitions of Terms Specific to This Standard:
3.2.1 lot, n—The term “lot” as used in this specification is defined as follows:

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1 This specification is under the jurisdiction of ASTM Committee B02 on Nonferrous Metals and Alloys and is the direct responsibility of Subcommittee B02.02 on Refined Lead, Tin, Antimony, and Their Alloys.


2 For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For Annual Book of ASTM Standards volume information, refer to the standard’s Document Summary page on the ASTM website.


3.2.1.1 Discussion—For solid solder metal, a lot consists of all solder of the same type designation, produced from the same batch of raw materials under essentially the same conditions, and offered for inspection at one time.

3.2.1.2 Discussion—For flux–core solder, a lot consists of all solder of the same core mixture, produced from the same batch of raw materials under essentially the same conditions and offered for inspection at one time.

3.2.2 lot number, n—The term “lot number” as used in this specification refers to an alphanumeric or numerical designation for a lot which is traceable to a date of manufacture.

4. Classification

4.1 Type Designation—The type designation uses the following symbols to properly identify the material:

4.1.1 Alloy Composition—The composition is identified by a two-letter symbol and a number. The letters typically indicate the chemical symbol for the critical element in the solder and the number indicates the nominal percentage, by weight, of the critical element in the solder. The designation followed by the letters A or B distinguishes between different alloy grades of similar composition (see Table 1).

4.1.2 Form—The form is indicated by a single letter in accordance with Table 2.

4.1.3 Flux Type—The flux type is indicated by a letter or combination of letters in accordance with Table 3.

4.1.4 Core Condition and Flux Percentage (applicable only to flux-cored solder)—The core condition and flux percentage is identified by a single letter and a number in accordance with Table 4.

4.1.5 Powder Mesh Size and Flux Percentage (applicable only to solder paste)—The powder mesh size and flux percentage is identified by a single letter and a number in accordance with Table 5.

5. Ordering Information

5.1 Orders for material under this specification indicate the following information, as required, to adequately describe the desired material.

5.1.1 Type designation (see 4.1),
5.1.2 Detailed requirements for special forms,
5.1.3 Dimensions of ribbon and wire solder (see 9.2),
5.1.4 Unit weight,
5.1.5 Packaging (see Section 18),
5.1.6 Marking (see Section 17),
5.1.7 ASTM specification number and issue, marked on (a) purchase order and (b) package or spool, and
5.1.8 Special requirements, as agreed upon between supplier and purchaser.

6. Materials and Manufacture

6.1 The producer must have each lot of solder metal as uniform in quality as practicable and of satisfactory appearance in accordance with best industrial practices. Each bar, ingot, or other form in which the solder is sold must be uniform in composition with the entire lot.

7. Chemical Composition

7.1 Solder Alloy—The solder alloy composition is as specified in Table 1.

Note 1—By mutual agreement between supplier and purchaser, analysis may be required and limits established for elements or compounds not specified in Table 1.

7.2 Flux (applicable to flux-core ribbon, wire, and solder paste):

7.2.1 Type R—The flux is composed of Grade WW or WG gum rosin of Test Methods D509. The rosin shall have a toluene–insoluble matter content of not more than 0.05 weight % in accordance with Test Method D269, a minimum acid number of 160 mg KOH/1 g sample in accordance with Test Methods D465, and a minimum softening point of 70°C in accordance with Test Methods E28, and a minimum saponification number of 166 in accordance with Test Methods D464.

When solvents or plasticizers are added, they must be nonchlorinated.

7.2.2 Type RMA—The flux is composed of rosin conforming to 7.2.1. Incorporated additives provide a material meeting the requirements of 8.1.2 for type RMA. When solvents or plasticizers are added, they must be nonchlorinated.

7.2.3 Type RA—The flux is composed of rosin conforming to 7.2.1. Incorporated additives provide a material meeting the requirements of 8.1.2 for type RA. When solvents or plasticizers are added, they must be nonchlorinated.

7.2.4 Type OA—The flux is composed of one or more water-soluble organic materials.

7.2.5 Type OS—The flux is composed of one or more water-insoluble organic materials, other than Types R, RMA, and RA, which are soluble in organic solvents.

7.2.6 Type IS—The flux is composed of one or more inorganic salts or acids with or without an organic binder and solvents.

8. Physical Properties and Performance Requirements

8.1 Solder Paste—Solder paste must exhibit smoothness of texture (no lumps) and the absence of caking and drying.

8.1.1 Powder Mesh Size—The solder powder mesh size shall be as specified (see 5.1.1 and 4.1.5) when the extracted solder powder is tested as specified in 13.4.

8.1.2 Viscosity—The viscosity of solder paste and the method used to determine the viscosity must be agreed upon between the supplier and purchaser. The following variables must be taken into account when relating one viscosity measurement to another type of viscometer used, spindle size and shape, speed (r/min), temperature of sample, and the use or non-use of a helipath.

8.2 Requirements for Flux—The flux must meet the physical and performance requirements specified in Table 6 as applicable.

8.2.1 Solder Pool—When solder is tested as specified in 13.3.2, there must be no spattering, as indicated by the presence of flux particles outside the main pool of residue. The flux must promote spreading of the molten solder over the coupon to form integrally thereon a coat of solder that shall
<table>
<thead>
<tr>
<th>Composition, %</th>
<th>Melting Range</th>
<th>UNS Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alloy Grade</td>
<td>Solidus °F</td>
<td>Liquidus °F</td>
</tr>
<tr>
<td>Sn96</td>
<td>430</td>
<td>221</td>
</tr>
<tr>
<td>Sn95</td>
<td>430</td>
<td>221</td>
</tr>
<tr>
<td>Sn94</td>
<td>430</td>
<td>221</td>
</tr>
<tr>
<td>St65</td>
<td>450</td>
<td>233</td>
</tr>
<tr>
<td>E10</td>
<td>440</td>
<td>225</td>
</tr>
<tr>
<td>HA</td>
<td>420</td>
<td>216</td>
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<tr>
<td>HB</td>
<td>460</td>
<td>238</td>
</tr>
<tr>
<td>HA</td>
<td>440</td>
<td>225</td>
</tr>
<tr>
<td>PT</td>
<td>430</td>
<td>221</td>
</tr>
<tr>
<td>AC</td>
<td>403</td>
<td>206</td>
</tr>
<tr>
<td>OAm</td>
<td>420</td>
<td>216</td>
</tr>
<tr>
<td>AM</td>
<td>430</td>
<td>220</td>
</tr>
<tr>
<td>TC</td>
<td>440</td>
<td>225</td>
</tr>
<tr>
<td>WS</td>
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<td>225</td>
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<tr>
<td>Sn70</td>
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<td>183</td>
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<td>Sn63</td>
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<td>183</td>
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<td>Sn62</td>
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<td>179</td>
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<td>Sn60</td>
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<td>Sn40</td>
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<td>Sn2</td>
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<td>185</td>
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<tr>
<td>Ag1.5</td>
<td>588</td>
<td>308</td>
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<tr>
<td>Ag2.5</td>
<td>580</td>
<td>308</td>
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<tr>
<td>Ag3.5</td>
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<tr>
<td>Ag4.5</td>
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<td>308</td>
</tr>
<tr>
<td>Ag5.5</td>
<td>580</td>
<td>308</td>
</tr>
</tbody>
</table>

For purposes of determining conformance to these limits, an observed value or calculated value obtained from analysis shall be rounded to the nearest unit in the last right-hand place of figures used in expressing the specified limit, in accordance with the rounding method of Practice E29.

Grades E and OA are covered by U.S. patents held by Engelhard Corp, Mansfield, MA, and Oatey Co. Cleveland, OH respectively. Federated Fry Metals, Altona, PA and Taracorp Inc., Atlanta, GA have applied for patents on grades AC and TC respectively.

OA value for Fe 10 was corrected editorially.

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feather out to a thin edge. The complete edge of the solder pool must be clearly visible through the flux residue.

8.2.2 Dryness—When solder is tested as specified in 13.3.2, the surface of the residue must be free of tackiness, permitting easy and complete removal of applied powdered chalk.

8.2.3 Chlorides and Bromides Test—When the extracted flux is tested as specified in 13.3.6, the test paper will show no chlorides or bromides by a color change of the paper to off-white or yellow white.

8.2.4 Copper Mirror Test—When tested as specified in 13.3.7, the extracted flux will have failed the test if, when examined against a white background, complete removal of the copper film is noted, as evidenced by the white background showing through, and must be rejected. Discoloration of the copper due to a superficial reaction or to only a partial reduction of the thickness of the copper film is not cause for rejection.

9. Dimensions and Unit Weight

9.1 Bar and Ingot Solder—The dimensions and unit weight of bar and ingot solder will be as agreed upon between supplier and purchaser.

9.2 Wire solder (solid and flux-cored)—The dimensions and unit weight of wire solder are specified in 5.1.3 and 5.1.4. The tolerance on the specified outside diameter shall be ±5% or ±0.002 in. (0.05 mm), whichever is greater.

9.3 Other Forms:
9.3.1 Dimensions for ribbon and special forms will be agreed upon between supplier and purchaser.
9.3.2 The unit weight of solder paste is specified in 5.1.4.

10. Workmanship, Finish, and Appearance

10.1 All forms of solder must be processed in such a manner as to be uniform in quality and free of defects that will affect life, serviceability, or appearance.

11. Sampling

11.1 Care must be taken to ensure that the sample selected for testing is representative of the material. The method of sampling consists of one of the following methods:
11.1.1 Samples taken from the final solidified cast or fabricated product.
11.1.2 Representative samples obtained from the lot of molten metal during casting. The molten sample is poured into a cool mold, forming a bar approximately 1/4 in. (6.4 mm) thick.

11.2 Frequency of Sampling—Frequency of sampling for determination of chemical composition shall be in accordance with Table 7. For spools and coils, the sample is obtained by cutting back 6 ft (1.8 m) of wire from the free end and then taking the next 6 ft for test. In other forms, an equivalent sample is selected at random from the container.
11.3 Other Aspects of Sampling—Other aspects of sampling conforms in the case of bar and ingots, to Practice E88. For fabricated solders the appropriate reference is Practice E55.

12. Specimen Preparation

12.1 Flux-Cored Ribbon and Wire Solder and Solder Paste—Each sample of flux-cored ribbon or wire solder or solder paste is melted in a clean container under oil and mixed thoroughly. After the flux has risen to the top, the alloy is poured carefully into a cool mold (care should be taken to allow the flux and alloy to separate completely), forming a bar approximately 1/4 in. (6.4 mm) thick.

12.1.1 Flux Extraction Procedure:
12.1.1.1 Flux-Cored Solder—The flux core is extracted as follows: Cut a length of the flux-cored solder weighing approximately 150 g and seal the ends. Wipe the surface clean with a cloth moistened with acetone. Place the sample in a
beaker, add sufficient distilled water to cover the sample, and boil for 5 to 6 min. Rinse the sample with acetone and allow to dry. Protecting the solder surface from contamination, cut the sample into ¼ in. (9.5 mm) (maximum) lengths without crimping the cut ends. Place the cut lengths in an extraction tube of a chemically clean soxhlet extraction apparatus and extract the flux with reagent grade, 99% isopropyl alcohol until the return condensate is clear. The resistivity of water extract, copper mirror, and chlorides and bromides tests are performed using a test solution prepared by concentrating the solids content of the test solution to approximately 35% by weight by evaporation of the excess solvent. The exact solids content of the test solution are determined on an aliquot, dried to constant weight in a circulating air oven maintained at 85 ± 3°C.

12.1.1.2 Solder Paste—The flux is extracted as follows: Place 200 mL of reagent grade, 99% isopropyl alcohol in a chemically clean Erlenmeyer flask. Add 40 ± 2 g of solder paste to the flask, cover with a watch glass, and boil for 10 to 15 min using medium heat. Allow the powder to settle for 2 to 3 min and decant the hot solution into a funnel containing filter paper, collecting the flux extract in a chemically clean vessel.

Note 2—The solution in isopropyl alcohol does not necessarily have to be clear. The resistivity of water extract and chlorides and bromides tests shall be performed using a test solution prepared by concentrating the solids content in the flux extract solution to approximately 35% by weight by evaporation of the excess solvent. The exact solids content of the test solution shall be determined on an aliquot, dried to constant weight in a circulating air oven maintained at 85 ± 3°C.

12.2 Solid Ribbon and Wire Solder—Each sample of solid ribbon and wire solder is prepared in accordance with 12.1, as applicable.

12.3 Bar and Ingot Solder—Each sample piece is cut in half and one half marked and held in reserve. The remaining half is melted in a clean container, mixed thoroughly and poured into a cool mold, forming a bar approximately ¼ in. (6.4 mm) thick. Sampling is performed by one of the following methods:

12.3.1 Sawing—Saw cuts are made across the bar at equal intervals of not more than 1 in. (2.5 cm) throughout its length. If it is impractical to melt the bar or ingot as specified above, saw cuts are made across each piece at equal intervals of not more than 1 in. (2.5 cm) throughout its length. No lubricants are used during sawing. The specimen consists of not less than 5 oz (143 g) of mixed sawings.

12.3.2 Drilling—The bar is drilled at least halfway through from two opposite sides. A drill of about ½ in. (12.7 mm) in diameter is preferred. In drilling, the holes are placed along a diagonal line from one corner of the pig to the other. The drillings are clipped into pieces not over ½ in. (12.7 mm) in length and mixed thoroughly. The specimen consists of not less than 5 oz (143 g).

13. Test Methods

13.1 Visual and Dimensional Examination:

13.1.1 Ribbon and Wire Solder (Solid and Flux-Cored)—Ribbon and wire solder must be examined to verify that the dimensions, unit weight, and workmanship are in accordance with the applicable requirements.

13.1.2 Solder Paste—Solder paste must be examined for smoothness of texture (no lumps), caking, drying, unit weight, and workmanship in accordance with the applicable requirements.

13.1.3 Bar and Ingot Solder—Bar and ingot solder must be examined to verify that the unit weight, marking, and workmanship are in accordance with the applicable requirements.

13.2 Alloy Composition—In case of dispute, the chemical analysis is made in accordance with Test Methods E46, Method E51, and Methods E87.

13.3 Flux:

13.3.1 Determination of Weight Percent of Flux:

13.3.1.1 Select a minimum of 20 g of flux-core ribbon or wire or solder paste. Weigh the sample in a clean porcelain crucible determining the weight to the nearest 0.01 g. Heat until the solder is completely molten. Carefully stir the molten solder a few times to free any entrapped flux. Allow the solder to cool until it solidifies; clean thoroughly of flux residues and reweigh the solder.

13.3.1.2 Calculation—Calculate the weight percent of flux as follows:

### TABLE 6 Requirements for Flux

<table>
<thead>
<tr>
<th>Test</th>
<th>Type R</th>
<th>Type RMA</th>
<th>Type RA</th>
<th>Other Flux Types</th>
<th>Method Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight of flux</td>
<td>see Table 4</td>
<td>see Table 4</td>
<td>see Table 4</td>
<td>see Table 4</td>
<td>13.3.1</td>
</tr>
<tr>
<td>Solder pool</td>
<td>see 8.2.1</td>
<td>see 8.2.1</td>
<td>see 8.2.1</td>
<td>see 8.2.1</td>
<td>13.3.2</td>
</tr>
<tr>
<td>Spread factor</td>
<td>80 min</td>
<td>80 min</td>
<td>80 min</td>
<td>not required</td>
<td>13.3.3</td>
</tr>
<tr>
<td>Dryness</td>
<td>see 8.2.2</td>
<td>see 8.2.2</td>
<td>see 8.2.2</td>
<td>not required</td>
<td>13.3.4</td>
</tr>
<tr>
<td>Resistivity of water extract (1-1 cm)</td>
<td>100 000 min</td>
<td>100 000</td>
<td>50 000</td>
<td>not required</td>
<td>13.3.5</td>
</tr>
<tr>
<td>Chlorides and bromides</td>
<td>see 8.2.3</td>
<td>see 8.2.3</td>
<td>not required</td>
<td>not required</td>
<td>13.3.6</td>
</tr>
<tr>
<td>Copper mirror</td>
<td>pass</td>
<td>pass</td>
<td>not required</td>
<td>not required</td>
<td>13.3.7</td>
</tr>
</tbody>
</table>

A Applicable only to composition 60/40.
B Applicable only to composition 60/40 in the form of flux-core wire or solderpaste.
C Applicable only to composition 60/40 in the form of flux-core wire.
D Applicable only to flux-core wire and solderpaste.

### TABLE 7 Frequency of Sampling

<table>
<thead>
<tr>
<th>Size of Lot, lb (kg)</th>
<th>Number of Samples (spools, coils, containers or pieces)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 1000 (450), incl</td>
<td>3</td>
</tr>
<tr>
<td>Over 1000 to 10 000 (450 to 4500), incl</td>
<td>5</td>
</tr>
<tr>
<td>Over 10 000 (4500)</td>
<td>10</td>
</tr>
</tbody>
</table>
13.3.3.2 Solder Pool (applicable only to composition 60/40)—For each sample being tested, three coupons 1.5 in. (38 mm) square shall be cut from 0.063 in. (1.6 mm) thick sheet copper. For flux Type IA only, the coupons shall be cut from cold-rolled commercial sheet steel, approximately 0.063 in. thick. The coupons are degreased by immersion in trichloroethylene or other suitable short-chain solvent. Both surfaces of cold-rolled commercial sheet steel, approximately 0.063 in. thick, must not exceed that of a 0.375 in. (9.5 mm) diameter sphere. The coupons are cleaned to a bright finish, using a 10 % fluoroboric acid dip. One corner of each coupon is bent upwards to permit evidence of spattering of flux.

The coupons in the oven, the flux residue is not removed. The coupons are allowed to cool for 1/2 h. Powdered chalk is dusted onto the surface of the residual flux and the ability to remove the chalk from the surface of the flux by light brushing is observed.

13.3.3.3 Calculation—The loss in weight of the solder slab in water is divided by five. This is the volume, $V$, of the solder to the nearest 0.001 cm$^3$. The diameter, $D$, of the equivalent sphere is $1.2407\sqrt[3]{V}$. The spread factor is calculated in accordance with the following formula:

$$ F = \frac{C - S}{C} \times 100 $$

where:

$F$ = weight percent of flux,
$C$ = initial weight of solder sample, g, and
$S$ = final weight of solder sample, g.

13.3.3.4 Dryness (applicable only to composition Sn60, flux Types R, RMA, and RA in the form of flux-core wire or solder paste)—The dryness test is performed on samples prepared in accordance with 13.3.3.1 and 13.3.3.2(a) except that after heating the coupons in the oven, the flux residue is not removed. The coupons are allowed to cool for 1/2 h. Powdered chalk is dusted onto the surface of the residual flux and the ability to remove the chalk from the surface of the flux by light brushing is observed.

13.3.5 Resistivity of Water Extract (applicable only to flux Types R, RMA, and RA)—The resistivity of water extract is determined using the flux test solution. Five watch glasses and five acid/alkali resistant, tall form graduated beakers are used. Each beaker is filled to the 50 mL mark with distilled water. The beakers are covered with watch glasses to protect the contents from contamination. The beakers’ dimensions are such that when the conductivity cell is immersed in 50 mL of liquid contained therein, the electrodes are fully covered. Each cleaned beaker is filled to the 50 mL mark with distilled water. The beakers are immersed in a water bath maintained at 23 ± 2°C. When thermal equilibrium is reached, the resistivity of the distilled water in each beaker is measured at this temperature with a conductivity bridge using a conductivity cell with a cell constant of approximately 0.1. The resistivity of the distilled water in each beaker must not be less than 500 000 $\Omega \cdot$ cm. If the resistivity of the water in any beaker is less than 500 000 $\Omega \cdot$ cm, the complete process above must be repeated. Two of these beakers are retained as controls. Add 0.100 ± 0.005 cm$^3$ of the flux test solution to each of the other three beakers by washing with alcohol, and the loss of weight in water determined to the nearest 0.001 g.
means of a calibrated dropper or microlitre syringe. The heating of all five beakers is started simultaneously. As the contents of each beaker comes to a boil, the boiling time is for 1 min followed by quick cooling of the beakers, to the touch, under running tap water or by immersing in ice water. The cooled, covered beakers are placed in a water bath maintained at 23 ± 2°C. When the thermal equilibrium is reached, the resistivity in each of the five beakers is determined at this temperature as follows:

13.3.5.1 Thoroughly wash the conductivity cell with distilled water and immerse it in the water extract of one sample. Make instrument reading.

13.3.5.2 Thoroughly wash conductivity cell in distilled water and immerse in a water control. Make instrument reading.

13.3.5.3 Thoroughly wash conductivity cell in distilled water and immerse in a water extract. Make instrument reading.

13.3.5.4 Thoroughly wash conductivity cell in distilled water and continue measuring resistivities of the remaining control and water extract.

13.3.5.5 The resistivity of each of the controls must not be less than 500 000 Ω-cm. If the control value is less than 500 000 Ω-cm, it indicates that the water was contaminated with water–soluble ionized materials and the entire test must be repeated. The mean of the specific resistivities of the water extracts of the flux must be calculated.

13.3.6 Chlorides and Bromides Test (applicable only to flux Types R and RMA)—One drop of the flux test solution (approximately 0.05 mL/drop) is placed on a small dry piece of silver chromate test paper. The drop shall remain on the test paper for 15 s prior to immersing the test paper in reagent grade 99 % isopropanol alcohol for 15 s to remove residual organic materials. The test paper is dried for 10 min. The test paper is visually examined for color change.

13.3.7 Copper Mirror Test (applicable only to flux Types R and RMA in the form of flux-core wire):

13.3.7.1 Preparation of the Control-Standard Flux—A control-standard flux is prepared by using 35 % weight of Grade WW gum rosin conforming to Test Methods D509 dissolved in reagent grade 99 % isopropanol alcohol.

13.3.7.2 Preparation of Copper Mirror—A copper mirror consists of a vacuum–deposited film of pure copper metal on one surface of a flat sheet of clear, polished glass. The thickness of the copper film must be uniform and must permit 10 ± 5 % transmission of normal incident light of 5000 A units as determined with any suitable standard photoelectric spectrophotometer. To prevent oxidation of the copper mirror, it is recommended that the mirrors be stored in closed containers which have been flushed with nitrogen. Immediately prior to testing, the copper mirror is immersed in a 5 % solution of ethylene diamine tetra acetic acid or similar chelating agent for copper oxide, rinsed thoroughly in running water, immersed in clean ethyl or methyl alcohol, and dried with clean, oil-free air. The copper film is examined in good light. The copper mirror is acceptable if no oxide film is visible and the copper film shows no visible damage.

13.3.7.3 Procedure—Approximately 0.05 mL of the flux test solution and 0.05 mL of the control–standard flux is placed adjacent to each other on the face of a flat, vacuum–deposited copper mirror. The dropper must not be permitted to touch the copper surface, and the mirror is protected at all times from dirt, dust, and fingerprints. The mirror is placed in a horizontal position at 23 ± 2°C and 50 ± 5 % relative humidity in a dust–free cabinet for 24 ± 1/2 h. At the end of the 24-h storage period, the test flux and the control standard flux are removed by immersing the copper mirror in clean isopropyl alcohol. The clean mirror is examined visually for compliance of the test flux and the control–standard flux with 8.1.2. If the control–standard flux does not comply with 8.1.2, the test must be repeated using a new copper mirror.

13.4 Powder Mesh Size (applicable only to solder paste)—Place 200 mL of reagent grade, 99 % isopropanol in a chemically clean beaker. Add 40 ± 2 g of solder paste to the beaker, cover with a watch glass and boil for 10 to 15 min using medium heat. Allow the powder to settle for 2 to 3 min and decant the hot solution. Wash the powder with isopropanol until all of the flux is removed. Replace isopropanol with deionized or distilled water for solder pastes containing a water–soluble flux base. Completely dry the solder powder at 110°C (230°F) so that all particles are separated. A minimum of 80 % of the powder must pass through the appropriate size sieve (see 4.1.5) in order to be classified for that mesh size.

14. Inspection

14.1 Unless otherwise specified in the contract or purchase order, the supplier is responsible for the performance of all inspection requirements as specified herein. Except as otherwise specified in the contract or order, the supplier may use his own or any other facilities suitable for the performance of the inspection requirements specified unless disapproved by the purchaser. The purchaser reserves the right to perform any of the inspections set forth in the specification where such inspections are deemed necessary to ensure supplies and services conform to prescribed requirements.

14.1.1 Test Equipment and Inspection Facilities—Test and measuring equipment and inspection facilities of sufficient accuracy, quality, and quantity to permit performance of the required inspection must be established and maintained by the supplier.

15. Rejection and Rehearing

15.1 Material that fails to conform to the requirements of this specification may be rejected. Rejection must be reported to the producer or supplier promptly and in writing. In case of dissatisfaction with the results of the test, the producer or supplier may make claim for a rehearing.

16. Certification

16.1 When specified in the purchase order or contract a producer’s or supplier’s certification must be furnished to the purchaser that the material was manufactured, sampled, tested, and inspected in accordance with this specification and has been found to meet the requirements. When specified in the purchase order or contract, a report of the test results must be furnished.
17. **Product Marking**

17.1 The producer’s name or trademark must be stamped or cast on each bar or ingot. The alloy grade designation or nominal composition, or both, must be stamped on each bar or ingot for identification along with the specification number.

17.2 Each spool or container must be marked to show the specification number, type designation, dimensions, and unit weight of wire or other form and lot number. The producer’s name or trademark must be marked on the spool or container.

18. **Packaging and Package Marking**

18.1 The material must be packaged to provide adequate protection during normal handling and transportation. The type of packaging and gross weight of containers will, unless otherwise agreed upon, be at the producer’s or supplier’s discretion, provided that they are such as to ensure acceptance by common or other carriers for safe transportation to the delivery point.

18.1.1 For bar and ingot solder a lot number must be marked on each shipping container or inside package.

18.1.2 When special preservation, packaging and packing requirements are agreed upon between purchaser and supplier, marking for shipment of such material must be in accordance with Fed. Std. No. 123 for civil agencies and MIL-STD-129 for military agencies.

18.2 Each shipping container must be marked with the purchase order number, unit weight, and producer’s name or trademark.

19. **Keywords**

19.1 bar; flux; flux cored solder; ingot; lead–silver alloys; lead–tin alloys; lead–tin–silver alloys; powder; ribbon; solder alloy; solder metal; solder uses; tin–antimony alloys; tin–copper alloys; tin–silver alloys; wire

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ANNEX

(Mandatory Information)

A1. INTENDED USE

A1.1 **Alloy Compositions:**

A1.1.1 Sn96—This is a special-purpose solder with a higher joint strength than tin–lead solders. It is intended for use in the food processing industry because of its nontoxic characteristic. It also provides a fairly good color match to stainless steel.

A1.1.2 Sn95, Sn94, E, AC, AM, and WS—These alloys are intended for use in soldering medical components and jewelry applications, for joining copper pipe and tube intended for potable water systems, and for applications in the food industry. These alloys display excellent wetting and exhibit relatively high heat resistance.

A1.1.3 Sn70—This is a special-purpose solder where a high tin content is necessary. It is often used for soldering zinc and for coating metals.

A1.1.4 Sn63—This tin–lead eutectic solder is commonly used for soldering printed circuit boards where temperature limitations are critical and in applications where an extremely short melting range is required.

A1.1.5 Sn62—This is a special-purpose solder widely used for soldering silver coated surfaces.

A1.1.6 Sn60—Similar to Sn63, this solder is preferred for soldering electrical and electronic connections and for coating metals.

A1.1.7 Sn50—This general purpose alloy can be used for non-critical electrical soldering and applications such as joining sheet metal, pipe, tubing and other structural shapes.

A1.1.8 Sn45—This is a general purpose alloy similar to Sn50.

A1.1.9 Sn40A—This alloy can be used for the same purposes as alloy Sn50, but it is not as workable in bit soldering or sweating. It is frequently used for dip soldering and as a wiping solder for joining lead pipes and cable sheaths.

A1.1.10 Sn40B—This alloy is similar to Sn40A, but it is not recommended for use on galvanized iron.

A1.1.11 Sn35A—This is a plumber’s solder similar to alloy Sn35B but with a lower antimony content.

A1.1.12 Sn35B—This is the customary wiping or plumber’s solder. Higher antimony content in wiping solders promotes fine grain size and greater strength.

A1.1.13 Sn30A—This alloy is used as an automobile–body solder and for removing heat–strippable insulation during high temperature (700 to 900°F) tinning of wires.

A1.1.14 Sn30B—This alloy is used as an automobile–body solder for filling dents and seams.

A1.1.15 Sn25A and Sn25B—These alloys are for uses similar to that for alloy Grades Sn20 and Sn30.

A1.1.16 Sn20A—This is an automobile–body solder with a lower antimony content than alloy Sn20B.

A1.1.17 Sn20B—This is widely used automobile–body solder for filling dents and seams, and for general purposes such as protective coatings on steel sheet where a high tin content alloy is not required.

A1.1.18 Sn15—This alloy is used for coating and joining metals.
A1.1.19 Sn10A—This alloy is used for coating and joining metals, and where soldered connections will be exposed to high operating temperatures exceeding 400°F (204°C).

A1.1.20 Sn10B—Similar to Sn10A, this alloy minimizes the leaching of silver from silver alloy coated surfaces. It is used in hybrid microelectronic and automotive electronic applications where a high service temperature is encountered.

A1.1.21 Sn5—This alloy is used for coating and joining metals, and where soldered connections will be exposed to high operating temperatures exceeding 475°F (246°C). Its wetting ability is not as good as Sn10A.

A1.1.22 Sn2—This alloy has been used to solder automobile radiator cores.

A1.1.23 Sb5—This alloy is used for electrical and electronic connections subjected to peak temperatures of approximately 465°F. It is also used for sweating of copper tubing in solar heating, plumbing and refrigeration equipment.

A1.1.24 Ag1.5—This alloy is used interchangeably with alloy Ag2.5, but has a better shelf life and does not develop a black surface deposit when stored under humid environmental conditions.

A1.1.25 Ag2.5—The alloy is for use on copper, brass and similar metals with torch heating. It requires the use of a flux having a zinc chloride base to produce a good joint on untinned surfaces. A rosin flux is unsatisfactory on untinned surfaces. This alloy is susceptible to corrosion under humid environmental conditions.

A1.1.26 Ag5.5—This alloy will develop a shearing strength of 1500 psi at 350°F (177°C). When soldering hard–drawn brass or copper, the application temperature should not exceed 850°F (454°C). A typical application is on thermocouples for aircraft engines where relatively high operating temperatures will not affect strength of the solder. Precautions noted for Ag2.5 also apply.

A1.1.27 HA, PT, and OA—A lead-free solder for joining copper plumbing systems. It has a lower melting temperature than Grade Sb5 and is suitable for filling connections with wider clearances.

A1.1.28 HN, HB and TC—A lead-free solder for joining copper plumbing systems. This solder has a wide liquidus/solidus range making it useful for filling solder connections that have wide clearances. It can also be used where service conditions require a solder with a higher melting temperature.

A1.2 Soldering of Zinc and Cadmium—In as much as zinc and cadmium appear to form intermetallic alloys with the antimony in the solder, compositions Sn40B, Sn35B, Sn30B, Sn25B, Sn20B and Sb5 should not be used for soldering zinc or cadmium, or zinc–coated or cadmium-coated iron or steel. These intermetallic alloys have high melting points which inhibit the flow of the solder, resulting in brittle joints.

A1.3 Flux Type:

A1.3.1 Type R—Type R is intended for use in the preparation of soldered joints for high reliability electrical and electronic applications.

A1.3.2 Type RMA—Type RMA provides a slightly more active fluxing action than Type R and is intended for similar uses.

A1.3.3 Type RA—Type RA provides more active fluxing action than Type RMA. It should be used for soldering joints which are readily accessible so that the residues can be removed by cleaning agents and procedures commonly used in industry. Since the fumes and particulates given off during soldering may be corrosive and contaminate the area surrounding the joint, this too must be susceptible to effective cleaning by the combination of materials and procedures to be used. There are many standard electrical soldering applications that use this type of flux.

A1.3.4 Type OA—Type OA is used for general soldering purposes on copper, nickel, brass, etc. Some fluxes of this type can be used for electrical and electronic soldering applications but complete removal of flux is necessary after soldering to prevent corrosion and current leakage.

A1.3.5 Type OS—Type OS has uses similar to Type OA.

A1.3.6 Type IS—Type IS is intended for use, exclusive of that in electrical or electronic circuits, in the preparation of mechanical and structural joints for all solderable metals, other than aluminum, magnesium and their alloys.

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