



## Standard Guide for Preparation of Metal Surfaces for Adhesive Bonding<sup>1</sup>

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

### INTRODUCTION

It is impossible and impractical to present a single surface-preparation method for a metal group, such as aluminum alloys, stainless steels, magnesium alloys, etc. The wide variety of individual alloys and heat treatments under each group, the fact that certain adhesives exhibit specific compatibilities with surface preparation and the complexity and nature of parts being bonded preclude the use of an all-inclusive procedure for a metal group. Procedures for aluminum alloys are well standardized, possibly because more bonding has been done with these alloys. Preliminary tests should be conducted with the specific adhesive and the exact lot of metals to determine performance. This is especially true for stainless steel. The adhesive manufacturers' recommendation should also be considered.

### 1. Scope

1.1 This guide covers procedures that have proved satisfactory for preparing various wrought metal surfaces for adhesive bonding. It does not address the preparation of cast metals.

1.2 The values stated in SI units are to be regarded as the standard. The values given in parentheses are for information only.

1.3 Surface preparation methods are included for:

| Metal                    | Section |
|--------------------------|---------|
| Aluminum Alloys          | 7       |
| Stainless Steel          | 8       |
| Carbon Steel             | 9       |
| Titanium Alloys          | 10      |
| Magnesium Alloys         | 11      |
| Copper and Copper Alloys | 12      |

1.4 Procedures included herein are based on the commercial practice of numerous agencies and organizations. The methods may be revised or supplemented, as necessary, to include methods based on proven performance.

1.5 The surface preparation of metal systems used for qualification and quality control testing of the adhesive should be agreed upon by both manufacturer and user.

1.6 *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 establish appropriate safety and health practices and determine the applica-*

*bility of regulatory limitations prior to use.*<sup>2</sup>

### 2. Referenced Documents

#### 2.1 ASTM Standards:

D 907 Terminology Relating to Adhesives<sup>3</sup>

D 1125 Test Methods for Electrical Conductivity and Resistivity of Water<sup>4</sup>

D 2674 Methods of Analysis of Sulfochromate Etch Solution Used in Surface Preparation of Aluminum<sup>3</sup>

D 3933 Guide for Preparation of Aluminum Surfaces for Structural Adhesives Bonding (Phosphoric Acid Anodizing)<sup>3</sup>

#### 2.2 ASM International Document:<sup>5</sup>

Metals Handbook Volume V

#### 2.3 Military Specifications:<sup>6</sup>

MIL-A-8625 Type II, Anodic Coatings for Aluminum and Aluminum Alloys

MIL-M-3171 Magnesium Alloy, Processes for Pretreatment and Prevention of Corrosion

MIL-M-45202 Magnesium Alloy, Anodic Treatment of

#### 2.4 SAE Standards:<sup>7</sup>

Aerospace Material Specification 2470—Anodic Treatment of Aluminum Base Alloy, Chromic Acid Treatment

<sup>2</sup> A surfactant that has been found suitable for this purpose consists of a 50 % mixture of a cationic and a fluochemical surfactant such as Zonyl FSC manufactured by E.I. DuPont, Wilmington, DE 19898.

<sup>3</sup> *Annual Book of ASTM Standards*, Vol 15.06.

<sup>4</sup> *Annual Book of ASTM Standards*, Vol 11.01.

<sup>5</sup> Available from ASM International, Metals Park, OH 44073.

<sup>6</sup> Available from Standardization Documents, Order Desk, Bldg. 4, Section D, 700 Robbins Ave., Philadelphia, PA 19111, Attn: NPODS.

<sup>7</sup> Available from the Society of Automotive Engineers (SAE), 400 Commonwealth Dr., Warrendale, PA 15096.

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### 3. Terminology

3.1 *Definitions*—Many of the terms in this guide are defined in Terminology D 907.

### 4. Apparatus

4.1 *Trays or Pans*, suitable for preparation of laboratory specimens and small parts. Generally in production, a series of tanks containing the necessary solutions is arranged with overhead cranes to transport parts to be prepared. Tanks should be lined with suitable materials to resist the solutions, and methods for circulating the solutions and heating with temperature controls should be included if required.

4.2 *Hooks, Baskets, Clamps*, or other holding devices, may be used to convey parts being prepared. The use of a 300 Series stainless steel for such equipment is suggested. Other metals may be used if they are resistant to the solution used.

### 5. Conditions

5.1 *Handling*—All parts during and after processing and before bonding should be handled as little as possible and only with clean, lint-free cotton or nylon fabric gloves.

5.2 Water used for preparing solutions should be distilled, steam condensate, demineralized, deionized or otherwise treated, if necessary, to obtain a condition of not more than 50 ppm of solids and have a pH between 5.5 and 10, or not more than 50 ppm of dissolved solids and not over 30 micromhos conductance (see Test Methods D 1125). Rinsing may be done by spray or by dipping in a tank in which the water is circulated and constantly being renewed by an overflow method. After removing from any rinse the water-break test is commonly used. If the water film is discontinuous, it indicates that the process is unsatisfactory and must be repeated (see 5.5.4). Other methods, such as indicator paper and contact angle, have been used to determine the effectiveness of the process. Rinsing should be sufficiently long and vigorous to assure removal of soluble residue chemicals, or particles. Critical specific temperatures are specified for the rinse water in some processes. Where not specified, temperature is not critical. The number of times a process may be repeated depends upon the amount of metal removed in relation to the tolerance requirements for the part and whether clad or bare metal is used. Generally, not more than two repeat treatments are permitted.

NOTE 1—No entirely foolproof method exists to determine bondability of a metal surface after preparation. Trial bonding followed by testing and evaluation against mechanical strength standards should be conducted to determine initial and continuous effectiveness of a method.

5.3 *Solutions*—The life of solutions should be standardized. The useful life of solutions depends upon the number and size of the parts being prepared. Solutions should be sampled periodically and analyzed for materials pertinent to the particular treatment method, such as, titration for hexavalent chromium ( $\text{CrO}_3$ ), iron, chlorides, aluminum, etc. A pH reading or acid content of trichloroethane vapor degreasing solution, or both, should be obtained. Standards should be established and records kept for such tests on each lot or vat of solution. Records should indicate the continuous conditions of solutions, such as amount of constituents added to bring solutions within limits, amount of contaminants present and date prepared (see Methods D 2674).

5.4 *Room Conditions* for surface preparation as well as other adhesive bonding operations should be controlled for temperatures of 18 to 24°C (65 to 75°F), relative humidity of 40 to 65 %, air-filtered to remove dust and pressure maintained at slightly higher than ambient.

5.4.1 The time and conditions between surface preparation and applying adhesive primer, if used, and bonding are critical. Usually, the time should not exceed 8 h and parts should be covered or wrapped in Kraft paper. Prepared surfaces can change their characteristics on standing and adhesives vary widely in their tolerance of adherend surface conditions. In production bonding of complicated or large parts, the time interval may exceed 8 h. Investigation should be made to determine the time limitation of the adhesive and the rate of change on the surface of the adherend. On the basis of the results of such an investigation, it may be possible to extend the length of time and establish standards for the interval between surface preparation and adhesive application to fit production schedules. Contaminating operations in the area should be avoided. Especially detrimental are paint or other spraying operations, processes using powdered materials, oil vapors from pumps and other machinery and spraying of mold release agents.

#### 5.5 *Common Steps*:

5.5.1 Remove ink markings or stamped identifications from the metals. This can be accomplished by wiping with a cloth wetted with a variety of commercial solvents, such as acetone, methyl ethyl ketone, lacquer thinner and naphtha.

5.5.2 The preferred degreasing method is vapor degreasing with 1,1,1-trichloroethane. Commercial vapor degreasing equipment is usually used. This equipment has a deep tank with provision for heating a chlorinated solvent to 82 to 87°C (180 to 190°F). Parts are suspended in the vapor zone above the hot liquid and allowed to remain until there is condensation and run off (approximately 5 min). This step is repeated if necessary until all visible contamination is removed. The parts are raised above the tank and allowed to dry thoroughly. Previously bonded parts, such as honeycomb core materials, are usually not degreased due to difficulty in removing residue.

5.5.3 Following vapor degreasing, mechanical or chemical cleaning methods are usually employed.

5.5.4 A water-break test is a common method used to analyze surface cleanliness. This test depends on the observation that a clean surface (one that is chemically active or polar) will hold a continuous film of water, rather than a series of isolated droplets. This is known as a water-break-free condition. A break in the water film indicates a soiled or contaminated area. Distilled water should be used in the test, and a drainage time of about 30 s should be allowed. Any trace of residual cleaning solution should be removed or a false conclusion may be made. If a water-break-free condition is not observed on the treated surface, it should not be used for bonding. The surface should be recleaned until the test is passed. If continuous failures occur, the treating process itself should be analyzed to determine the cause of the problem.

### 6. Report

6.1 The report for all methods consists of flow charts, record sheets or other methods that indicate all steps in the surface

preparation used, with spaces for entries made upon completion of each step.

## ALUMINUM ALLOYS

### 7. Procedure

7.1 Aluminum alloys may be pretreated with many different methods, using both acidic and basic cleaners.

7.2 A typical pretreatment would normally involve the following steps:

7.2.1 Vapor degrease or non-etching alkaline cleaner,

7.2.2 Rinse,

7.2.3 Acid or alkaline, and

7.2.4 Rinse.

7.3 After 7.2.4, the parts may be dried or they may be given an anodizing treatment.

7.4 Typical pretreatments are listed in 7.4.1. Refer to the appropriate literature for details on cleaning techniques and procedures.

7.4.1 Alkaline degreasing solutions may be used instead of, or in addition to, vapor degreasing. Commercially available proprietary solutions may be used. A common degreasing method is to immerse for 10 min at 70 to 82°C (160 to 180°F) in the following solution by weight: 3.0 parts sodium metasilicate, 1.5 parts sodium hydroxide, 0.5 parts sodium dodecylbenzene sulfonate, such as Nacconol 90G<sup>8</sup>, to 133.0 parts water.

7.4.2 Acid etching solutions typically used are the following:

7.4.2.1 Sulfuric Acid/Sodium Dichromate (optimized FPL etch).

| Material                       | Concentration                          |
|--------------------------------|----------------------------------------|
| Sulfuric Acid (66°Be or Baume) | 38 to 41.5 oz/gal (287.9 to 310.0 g/L) |
| Sodium Dichromate              | 4.1 to 9.0 oz/gal (28–67.3 g/L)        |
| Aluminum Alloy—2024 bare       | 0.2 oz/gal (1.5 g/L), min              |
| Temperature                    | 66 to 71°C (150 to 160°F)              |
| Immersion Time                 | 12 to 15 min                           |

7.4.2.2 P-2 Etch (Sulfuric Acid/Ferric Sulfate).

| Material                     | Concentration                    |
|------------------------------|----------------------------------|
| Sulfuric acid (6.5 to 9.5 N) | 27 to 36 % by weight             |
| Ferric sulfate               | 135 to 165 g/L (18 to 22 oz/gal) |
| Temperature                  | 60 to 65°C (140 to 150°F)        |
| Immersion Time               | 10 to 12 min                     |

7.4.3 *Phosphoric Acid Anodizing* can be performed in accordance with Guide D 3933.

7.4.4 *Sulfuric Acid Anodizing*—A method found suitable for bare aluminum (nonclad), machined, or chemically milled parts which must be corrosion protected is to anodize with sulfuric acid. Do not seal the anodized parts with boiling water before bonding. Do not allow the time interval between treatment and bonding to exceed 72 h, and cover or wrap all parts during the interval with a non-contaminating vapor barrier material.

NOTE 2—Details of the process are contained in Specification AMS 2470E, or, Military Specification MIL-A8625.

7.4.5 *Proprietary Cleaners*—Some proprietary, commercially available metal surface preparations have been found

suitable for preparing the surfaces for adhesive bonding. Full-scale property tests should be run with the metal surfaces so prepared with the adhesive in question before accepting the process.

7.4.6 *Secondary Bonding*—When secondary bonding (bonding of parts that contain previously bonded areas) or repair of a disbonded assembly in service is necessary, it is usually impossible and undesirable to immerse these parts in surface-preparation solutions without destroying or harming the original adhesive bond. Satisfactory surface preparations have been accomplished by using paste-type etchants. These paste systems can be made by adding fumed silica to the sulfuric-dichromic acid solutions. The application of these paste systems allows for their use in fairly controlled areas. The etching time is generally 10 to 20 min. Exercise extreme care to ensure removal of all traces of the etchant. A thorough rinsing with distilled water may be necessary to ensure complete removal. Several commercial products are available. Some commercial compounds under the classification of “wash primers” have been found to perform satisfactorily in place of the paste type systems. Bond strengths obtained using either of these surface preparations are somewhat lower than those from the immersion processes.

7.4.7 *Brush Plate Bond Etch (Stylus Method)*:

7.4.7.1 *General*—This method of electrochemical surface preparation, although not widely known, has been effectively demonstrated for small batch runs where tanks and other capital-intensive methods might have been used. Almost all of the solutions, “brushes” (or styli), and electrical power supplies are proprietary. When this method is used, the entire “family” of proprietary items should be used; mixed usage of items between vendors should not occur.

7.4.7.2 *General Precleaning*—Removal of grease, oil, and other organic soils can be accomplished with virtually any solvent, including trichlorotrifluoroethane and similar non-flammable types.

7.4.7.3 *Mechanical Preparation*—Surfaces which are scaled, corroded, or otherwise oxidized should be abraded using a nonmetallic abrasive. Aluminum oxide-impregnated nylon matting, glass-bead blasting and aluminum oxide cloth are typical acceptable abrasives. Abrasive and debris should be wiped or water rinsed from the surface.

7.4.7.4 *Electrochemical Cleaning*—This is accomplished by immersing the wrapped electrode in an alkaline electrocleaner-type solution and swabbing the surface while applying a cathodic current at approximately 5 to 15 V. (The surface is the cathode.) Thorough rinsing follows.

7.4.7.5 *Anodic Etching*—This step is recommended for most wrought alloys of aluminum. The surface is anodically etched (again using a wrapped electrode) and a deep-etching solution. Five to fifteen volts are typical. Thorough rinsing should follow. The surface will exhibit a grey or lighter color.

7.4.7.6 *Bond Etching*—The solution of choice is applied using an anodic current, first at a low voltage and then at a higher value. Initial treatment is at about 5 to 10 V. Finishing voltages from 15 to 25 V are used. Terminal voltages and current decay schedules are typically specified. Thorough rinsing follows, usually in warm to hot running water.

<sup>8</sup> Nacconol 90G, available from the Stephan Co., Northfield, IL 60093, has been found suitable for this purpose.

7.4.7.7 *Final Rinse*—A final rinse or spray in a dilute chromic acid solution and another warm-to-hot water rinse.

7.4.7.8 *Drying*—The surface should be dried of water drops with dry oil-free compressed air or nitrogen and then left at ambient conditioning for about 1 h before the adhesive is applied. Longer dwell times should be avoided.

## STAINLESS STEEL

### 8. Procedure

8.1 *Mechanical Methods*—Mechanical methods such as wire brushing, rubbing with metal wool, hand sanding, dry sand blast, grit blast, or vapor honing may be used to prepare stainless steel. Exercise care to ensure that all abrasive materials (grit, sand, cloth, sand paper, or brushes) are free from contamination that may be spread or rubbed onto the steel. Dry grit or sand blasting tends to warp thin sheet materials; these methods are suited only for thick-section parts. Vapor blasting by water or steam and an abrasive is the most effective method and does not warp parts, if done carefully. Peening action of any blast process may work-harden the surface of some base metals. Hand sanding, rubbing with metal wool, and wire brushing are the least effective methods. Exercise care in using any mechanical method to prevent deep gouges or rough surfaces which are not conducive to good bonding. Wash, vaporize degrease, and brush or air blasted (dry oil-free air) surfaces to remove all traces of the abrasives. Apply the adhesive immediately after the treatment.

#### 8.2 Chemical Methods:

8.2.1 Adhesive primers or adhesives must be applied immediately after treatment and drying.

#### 8.2.2 Acid Etch (Sulfuric, Nitric, Hydrofluoric):

8.2.2.1 Etch for 10 min at 65 to 71°C (150 to 160°F) in the following solution by weight: 90 parts water, 37 parts sulfuric acid (sp gr 1.84), to 0.2 parts sodium dodecylbenzene sulfonate such as Nacconol 90G.

8.2.2.2 Rinse thoroughly and remove smut with a stiff brush if necessary.

8.2.2.3 Immerse for 10 min at room temperature in the following bright dip solution by weight: 88 parts water, 15 parts nitric acid (sp gr 1.42) to 2 parts hydrofluoric acid (35.35 %, sp gr 1.15).

8.2.3 *Acid Etch (Hydrochloric, Orthophosphoric, Hydrofluoric)*—Immerse for 2 min at approximately 93°C (200°F) in the following solution by weight heated by a boiling-water bath: 200 parts hydrochloric acid (sp gr 1.2), 30 parts orthophosphoric acid (sp gr 1.8), to 10 parts hydrofluoric acid (35.35 %, sp gr 1.15).

8.2.4 *Acid Etch (Sulfuric, Sodium Dichromate)*—Immerse for 15 min at 63 ± 3°C (155 ± 5°F) in the following solution by volume: 100 parts sulfuric acid (sp gr 1.84) to 30 parts saturated sodium dichromate solution.

8.2.5 *Sodium Metasilicate*—Immerse for 15 min at 63 ± 3°C (155 ± 5°F) in the following solution by weight: 47.2 parts water, 1.0 part sodium metasilicate to 1.8 parts of an anionic surfactant of the sodium salt, alkylaryl polyethersul-

fonate type (Triton X200<sup>9</sup> has been found suitable for this purpose).

#### 8.2.6 Acid Etch (Hydrochloric Sulfuric-Dichromate):

8.2.6.1 Immerse for 10 min at 60 to 65°C (140 to 150°F) in the following solution by weight: 45 parts water, 50 parts hydrochloric acid (sp gr 1.2), 10 parts formalin solution (40 %), to 2 parts hydrogen peroxide (concentration, 30 to 35 %).

8.2.6.2 Rinse thoroughly.

8.2.6.3 Immerse for 5 min at 50 to 65°C (140 to 150°F) in sulfuric acid-dichromate solution used for preparing aluminum (7.3.2.1).

8.2.6.4 Rinse thoroughly.

8.2.6.5 Dry at not over 93°C (200°F).

8.2.7 *Commercial Household Cleaner*—Some batches of stainless steel may be prepared by vigorous scouring with a wet cloth and a commercial household cleaner. This method should be used only in cases where other methods cannot be used. Somewhat lower bond strength results from this method compared to other methods described.

## CARBON STEEL

### 9. Procedure

9.1 *Mechanical Methods*—Carbon steel may be prepared using the same general methods as stainless steel. Sand blast, grit blast, or vapor hone may be used. Thin sheet steel may be warped due to peening of the blasting abrasives.

#### 9.2 Chemical Methods:

9.2.1 Thin parts and sheet materials may be prepared using methods for stainless steel. The major problem is in preventing oxidation of the surfaces. Oxidation can occur in a very short time, even before the parts are dry. Parts must be bonded immediately. The adhesive or adhesive primer must be applied immediately after drying.

#### 9.2.2 Nitric-Phosphoric Acid Etchant:

9.2.2.1 The nitric-phosphoric etchant is environmentally acceptable. It produces a smut-free surface that can be treated by other chemical processes.

9.2.2.2 Process parameters are as follows:

|                             |                   |
|-----------------------------|-------------------|
| Nitric acid—40° Baume       | 5 % by volume     |
| Phosphoric acid—85 % weight | 30 % by volume    |
| Surfactant <sup>9</sup>     | 0.01 % by volume  |
| Deionized water             | 64.99 % by volume |
| Temperature                 | 23°C (75°F)       |
| Immersion Time              | 5 min             |

## TITANIUM ALLOYS

### 10. Procedure

10.1 *Mechanical Abrasion*—See 8.1.

#### 10.2 Chemical Methods:

10.2.1 *Acid Etch (Hydrochloric, Orthophosphoric, Hydrofluoric):*

10.2.1.1 Immerse for 2 min at room temperature in the following solution: 841 mL orthophosphoric acid (reagent grade, 85 to 87 %), to 63 mL hydrofluoric acid (reagent grade, 60 %).

<sup>9</sup> Triton X200 is available from Rohm and Haas, Independence Mall West, Philadelphia, PA 19105.

### 10.2.1.2 Rinse.

10.2.1.3 Oven dry for 15 min at 88 to 93°C (180 to 200°F).

### 10.2.2 Acid Etch (Nitric Hydrofluoric):

10.2.2.1 Immerse for 15 min at 76°C (170°F) in the following solution: 6 to 8 fl oz of a caustic cleaner, such as Vitro-Klene<sup>10</sup>, or equivalent, to 3.6 L (1 gal) water.

10.2.2.2 Rinse in cold tap water.

10.2.2.3 Immerse for 5 min at room temperature in the following solution by weight: 48 % nitric acid (sp gr 1.5), 3 % ammonium bifluoride (technical), to 49 % water.

10.2.2.4 Rinse in cold tap water.

10.2.2.5 Air-dry at room temperature.

10.2.2.6 Immerse for 2 min at room temperature in the following solution: 50.0 g trisodium phosphate (technical), 8.9 g sodium fluoride (technical), 26.0 mL hydrofluoric acid (48 %), to water to make 3.6 L (1 gal) of solution.

10.2.2.7 Air-dry at room temperature.

10.2.3 *Stainless Steel Methods*—Processes used to prepare titanium alloys for adhesive bonding can be much the same as for stainless steel. Good bond strengths have resulted by bonding titanium alloys that have been anodized by proprietary processes.

10.2.4 *Chromic Acid/Fluoride Anodizing*—Stainless steel surface processes, as described in 8.2.4-8.2.6 have been found satisfactory.

## MAGNESIUM ALLOYS

### 11. Procedure

NOTE 3—The surface preparation methods for magnesium alloys are closely associated with corrosion prevention. The high reactivity of magnesium requires that corrosion-preventative coatings be applied for most service applications. Some of these coatings exhibit good surfaces for adhesive bonding. The major problem is to apply sufficient thickness of coating to prevent corrosion, but not so thick that the bond fails within the coating.

#### 11.1 Alkaline-Detergent Solution:

11.1.1 Degrease.

NOTE 4—**Caution** should be exercised during vapor degreasing of magnesium alloys. Contaminants, such as metal particles, oils, etc. may result in a fire or an explosion hazard.

11.1.2 Immerse for 10 min at 60 to 71°C (140 to 160°F) in an alkaline-detergent solution by weight: 95 parts water, 2.5 parts sodium metasilicate, 1.1 parts trisodium pyrophosphate, 1.1 parts sodium hydroxide, to 0.3 part sodium dodecylbenzene sulfonate such as Nacconol 90G.

11.1.3 Rinse thoroughly.

11.1.4 Dry at not over 60°C (140°F).

#### 11.2 Chromic Acid:

11.2.1 Degrease (see Note 4).

11.2.2 Immerse for 10 min at 71 to 88°C (160 to 190°F) in the following solution by weight: 4 parts water to 1 part chromic oxide (CrO<sub>3</sub>).

11.2.3 Rinse thoroughly.

11.2.4 Dry at not over 60°C (140°F).

NOTE 5—Methods 11.1 and 11.2 may be used as separate treatments; however, somewhat improved bond strengths can be obtained by using Method 11.1 followed by Method 11.2 with water rinse between the processes.

#### 11.3 Sodium Hydroxide-Chromic Acid:

11.3.1 Degrease (see Note 4).

11.3.2 Immerse for 5 to 10 min at 63 to 79°C (145 to 175°F) in the following solution by weight: 12 parts water, 1 part sodium hydroxide (commercial grade).

11.3.3 Rinse (water at or below room temperature).

11.3.4 Immerse for 5 to 15 min at room temperature in the following solution by weight: 123 parts water, 24 parts chromic oxide (CrO<sub>3</sub>) to 1.1 parts nitrate (commercial grade).

11.3.5 Rinse thoroughly.

11.3.6 Dry at not over 60°C (140°F).

11.4 *Anodic Treatment*—Light anodic treatments and various corrosion preventative treatments produce good surfaces for adhesive bonding.

NOTE 6—Details may be obtained from the magnesium alloy producers or from ASM Handbook, Vol V, and Military Specification MIL-M-45202, Type I, Classes 1, 2 and 3.

11.5 *Conversion Coatings and Wash Primers*—Some dichromate conversion coatings and wash primers designed for corrosion prevention are suitable for adhesive bonding. Preliminary tests should be conducted to determine suitability of these processes before acceptance.

NOTE 7—Details of the processes are found in ASM Metals Handbook, Vol V. Also, Military Specification MIL-M-3171 covers the methods.

## COPPER AND COPPER ALLOYS

### 12. Procedure

#### 12.1 Nitric Acid/Ferric Chloride:

12.1.1 Degrease.

12.1.2 Immerse for 1 to 2 min at room temperature in the following solution by weight: 197 parts water, 30 parts nitric acid (sp gr 1.42), to 15 parts ferric chloride solution (42 %).

12.1.3 Rinse thoroughly.

12.1.4 Dry as quickly as possible.

12.1.5 Apply adhesive immediately.

#### 12.2 Nitric Acid:

12.2.1 Bright dip in concentrated nitric acid at 16 to 21°C (60 to 70°F) for 15 s or until all corrosion has disappeared.

12.2.2 Rinse thoroughly.

12.2.3 Dry as fast as possible.

12.2.4 Apply adhesive immediately.

#### 12.3 Acid Etch (Sulfuric Acid-Dichromate-Ferric Sulfate):

12.3.1 Remove surface contamination by sanding, wire brushing, or sand blasting, if necessary.

12.3.2 Degrease.

12.3.3 Immerse for 10 min at 66°C (150°F) in the following solution by weight: 8 parts water, 1 part ferric sulfate (commercial), to 75 parts sulfuric acid (sp gr 1.84).

12.3.4 Rinse (water at or below room temperature).

12.3.5 Dry.

12.3.6 Immerse the parts until a bright clean surface has been obtained in the following solution by weight at room temperature: 17 parts water, 2 parts sulfuric acid (sp gr 1.84), to 1 part sodium dichromate.

<sup>10</sup> Vitro-Klene is available from Turco Purex Industries, P.O. Box 6200-T, Carson, CA 90749.

- 12.3.7 Rinse using cold tap water.
- 12.3.8 Dip in concentrated ammonium hydroxide.
- 12.3.9 Rinse in cold tap water.
- 12.3.10 Dry quickly.
- 12.3.11 Apply adhesive immediately.

12.4 *Chemical Treatment (Nitric Acid, Sodium Chlorite):*

NOTE 8—This method is suitable for copper alloys containing over 95 % copper. It is not suitable for adhesives containing chlorides or for hot-bonding polyethylene.

12.4.1 Degrease.

12.4.2 Immerse for 30 s at room temperature in the following solution by volume: 30 parts nitric acid (70 % technical) to 90 parts water.

12.4.3 Rinse in running cold tap water. Do not allow to dry.

12.4.4 Immerse for 2 to 3 min at 93 to 102°C (200 to 215°F) in the following solution: 4.01 oz sodium chlorite (technical), 1.34 oz trisodium phosphate, 0.67 oz sodium hydroxide, to 3.6 L (1.00 gal) water.

NOTE 9—If treating solution is to be used for a continuous operation or reused, it should be discarded after 4 to 6 h at the temperatures in 12.4.4.

12.4.5 Rinse thoroughly in running water until a neutral test is produced when touched with indicator paper.

12.4.6 Dry. Bond as soon as possible, but at least within the same working day.

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