

# METALLURGICAL ABSTRACTS

(GENERAL AND NON-FERROUS)

Volume 2

SEPTEMBER 1935

Part 9

## I.—PROPERTIES OF METALS

(Continued from pp. 365-370.)

\***The Softening of a Strained Metal [Aluminium] by Relaxation.** M. Kornfeld (*Physikal. Z. Sowjetunion*, 1934, 6, 329-342).—[In German.] The dependence of the velocity of softening of an aluminium single crystal on the initial strain and temperature of relaxation is investigated. The nature of the dependence of the flow limit on the relaxation temperature, established by K. in previous work (*Met. Abs.*, 1934, 1, 228, and this vol., p. 208) is now confirmed over a wide range of strains. Above 550° C. a second steady decrease of flow limit with decrease of relaxation temperature occurs, so that finally the original elastic characteristics of the crystal are recovered. A theory of relaxation is briefly discussed.—J. S. G. T.

\***Electrical Resistance of Pure Aluminium at Liquid Helium Temperatures.** H. A. Boorse and H. Niewodniczański (*Nature*, 1935, 135, 827-828).—Before becoming superconducting, the electrical resistance of very pure aluminium is constant between 4.2° and 2.2° abs.—E. S. H.

\***On the Supraconductivity of Aluminium.** W. H. Keesom (*Proc. K. Akad. Wet. Amsterdam*, 1933, 36, 381-383).—[In English.] See *Met. Abs.*, 1934, 1, 545.—L. A. O.

\***The Magnetostriction of Bismuth Single Crystals.** D. Shoenberg (*Proc. Roy. Soc.*, 1935, [A], 150, 619-637).—The transverse magnetostriction effect exhibited by bismuth single crystals in magnetic fields of strength below 20,000 oerstedt is investigated experimentally. The change of length, of the order  $5 \times 10^{-7}$  cm., was measured by a special magnification method based on a hydraulic principle. The results confirm the general features of the phenomenon observed by Kapitza in intense longitudinal magnetic fields and are expressed in terms of 8 magnetostriction moduli, values of which are given. The anomalous temperature variation of diamagnetic susceptibility of bismuth is discussed in connection with the theoretical significance of these moduli.—J. S. G. T.

\***Crystal Structure and Electrical Properties. V.—The [Electrical] Conductivity Surface of a Bismuth Crystal (Part 2).** O. Stierstadt (*Z. Physik*, 1935, 93, 676).—In continuation of previous work (*Met. Abs.*, 1934, 1, 77, 127, 484), the form of the electrical conductivity surface of a bismuth crystal in a transverse magnetic field is shown to be related, in general, to that characteristic of a longitudinal field. The surface, however, in the former case is not unique; characteristics of the surface are discussed.—J. S. G. T.

\***Properties of Calcium Obtained by Sublimation.** Paul Bastien (*Rev. Mét.*, 1935, 32, 120-136).—Very pure calcium (99.3%) is obtained by sublimation *in vacuo* and re-melting in argon. Observations on dilatation, differential thermoelectric properties, and hardness after quenching from various selected temperatures reveal 3 allotropic modifications of calcium. The transformation temperatures are 260° and 440° C. Dilatation curves of impure calcium show 2 supplementary anomalies which seem to be due to the action of atmospheric nitrogen on the warm metal. Sublimed calcium has a considerable capacity for deformation, which may be increased by hot-drawing at 420°-460° C. The impurities in industrial calcium reduce its capacity for

\* Denotes a paper describing the results of original research.

† Denotes a first-class critical review.

deformation to a very moderate value. Sublimed calcium shows a lower rate of corrosion in alcoholic hydrochloric acid than commercial varieties, the corrosion-resistance of which is decreased by the impurities present.—H. S.

**\*Investigations Relating to Heat Changes Associated with Metallic Transformations.** II.—[Cobalt]; III.—[Iron]. H. v. Steinwehr and A. Schulze (*Physikal. Z.*, 1935, **36**, 307-311, 419-423).—(II.—) The heat change associated with the  $\alpha$ - $\beta$  transformation of cobalt has been found to have the value  $0.1 \pm 0.03$  cal./gm.; that associated with the magnetic transformation has the value  $1.2 \pm 0.07$  cal./gm. (III.—) Values of the heat changes associated respectively with the magnetic ( $A_2$ ) and the  $A_3$  transformations of iron have been found to have the values ( $A_2$ )  $4.8 \pm 0.2$  cal./gm.; ( $A_3$ )  $6.2 \pm 0.08$  cal./gm.—J. S. G. T.

**\*The Electrical Resistance of Cobalt in a Longitudinal Magnetic Field.** Md. Sharf Alam (*Z. Physik*, 1935, **93**, 556-560).—The electrical resistance of a cobalt strip (purity 95%) in a longitudinal magnetic field is found to increase with increase of strength of field until a saturation value is attained, corresponding with a state of magnetic saturation of the metal, at a field-strength of about 1250 oerstedt. On subsequently decreasing and reversing the magnetic field the electrical resistance exhibits the phenomenon of hysteresis, the value of the resistance as the field is decreased being above that shown when the field is initially increased. The increase of resistance at saturation value is about 0.55-0.58% of the initial value of the resistance. A complete hysteresis cycle can be obtained by employing a cycle of values of the magnetizing field. In this cycle a field-strength of about 15 oerstedt is necessary to restore the resistance to its initial virginal value.—J. S. G. T.

**\*A Metallic Compound of Lithium and Ammonia. Electrical Conductivity and Galvanomagnetic Effects.** Hans Jaffe (*Z. Physik*, 1935, **93**, 741-761).—A saturated solution of lithium in liquid ammonia (containing about 4 mols. ammonia to 1 mol. lithium) freezes to form a solid compound at  $-180^\circ\text{C}$ .; the solid and liquid exhibit the phenomenon of metallic reflection, reflected light being copper-like in colour. The specific electric resistance of the saturated solution is about half that of mercury at room temperature and very nearly independent of temperature; the specific resistance of the solid compound at  $-190^\circ\text{C}$ . is reduced to about  $\frac{1}{4}$  and has a positive temperature coeff. of about 2%. The saturated solution of lithium in ammonia, with a density of 0.48, is the lightest of all liquids at room temperature. The solid compound shows a normal negative Hall effect; at  $-190^\circ\text{C}$ . the Hall coeff. has the value  $-1.38 \times 10^{-3}$  cm.<sup>3</sup>/amp. sec. It follows that exactly 1 free electron is present in the compound per atom of lithium. At  $-200^\circ\text{C}$ . a transformation of the compound occurs; this transformation is accompanied by a sudden fourfold increase in the value of the Hall coeff., and the marked dependence of the value of the electrical resistance of the compound on the strength of the magnetic field in which it is placed. The Hall coeff. for the saturated liquid is less than  $-2.5 \times 10^{-3}$  cm.<sup>3</sup>/amp. sec.—J. S. G. T.

**\*On the Recovery of Iron and Nickel from Cold-Work.** G. Tammann and G. Moritz (*Ann. Physik*, 1933, [v], **16**, 667-679).—The changes produced in the electrical resistance and temperature coeff. of resistance of iron and nickel by rolling, drawing, and annealing have been measured.—v. G.

**\*The Influence of Impurities in the Core-Metal on the Thermionic Emission from Oxide-Coated Nickel.** M. Benjamin (*Phil. Mag.*, 1935, [vii], **20**, 1-24).—The commoner metallic impurities, added in turn to a pure nickel oxide-coated core-metal, are found to have a very considerable effect on the thermionic emission. This phenomenon is explained on the assumption that the amount of barium metal in the coating is dependent on the reducing power of the impurity.—J. S. G. T.

**\*Oxidation of Silicon.** A. Sanfourche (*Rev. Mét.*, 1935, **32**, 182-188).—Silicon oxidizes at low temperatures, even at normal air temperature. As

with other oxidizable metals, the action is the more intense the finer the state of subdivision. The product of the oxidation is either silica or a lower oxide. The oxide film usually protects against further oxidation when it consists of silica. The action of hydrofluoric acid on the oxide film led Moissan and Siemens to assume the existence of an allotropic form of silicon, soluble in hydrofluoric acid. The hydrogen liberation does not correspond to dissolution of silicon, but is much less in amount and is attributed by S. to the action of a sub-oxide. The modification of silicon-aluminium alloys by alkaline salts is attributed to elimination of the oxide film from silicon, thereby permitting intimate mixing of the constituents.—H. S.

**\*The Rate of Polymorphic Transformations. New Investigations on Tin Pest.** Ernst Cohen and A. K. W. A. van Lieshout (*Z. physikal. Chem.*, 1935, [A], 173, 1-3).—The rate of transformation of white tin to grey and *vice versa*, is dependent on the number of transformations the metal has previously undergone, the medium (*e.g.* air or xylene) in which the transformation takes place, the previous thermal and mechanical history of the metal, and whether or not the metal is shaken during the change. Tin has been prepared in a form which is 7000 times as active as any previously known, so that it is possible to determine the rate of transformation at 1 atmosphere pressure in the neighbourhood of the transformation temperature; the rate is at 11.40° C. 6, 10.40° C. 12, and 9.40° C. 17.5 mm./hr. The observed phenomena are explained on the basis of the relation between the deformation and rate of transformation of enantiotropic metals.—B. Bl.

**\*Physico-Chemical Investigations on Tin.—The Temperature of Transformation White Tin  $\rightleftharpoons$  Grey Tin.** Ernst Cohen and A. K. W. A. van Lieshout (*Z. physikal. Chem.*, 1935, [A], 173, 32-34).—Previous determinations of the transformation temperature of tin were made only within a temperature interval of 2.3° C. Using the highly active tin prepared as described in an earlier paper (see preceding abstract), the transformation temperature was found to be  $13.2 \pm 0.1^\circ \text{C}$ .—B. Bl.

**\*The Rate of Polymorphic Transformations.—II.** Ernst Cohen and A. K. W. A. van Lieshout (*Z. physikal. Chem.*, 1935, [A], 173, 67-70).—The "region of indifference" observed by Bridgman and others in the transformation of polymorphic substance is not a specific property of the substance but is to be ascribed to the previous history of the specimen; by suitable pre-treatment this region can be caused to disappear.—B. Bl.

**\*The Rate of Polymorphic Transformations. III.—The Influence of Mechanical Deformation on the Rate of Transformation [of Tin].** Ernst Cohen, W. A. T. Cohen de Meester, and A. K. W. A. van Lieshout (*Z. physikal. Chem.*, 1935, [A], 173, 169-177).—Deformation of white tin by rolling, drawing, or bending accelerates very greatly the rate of transformation into grey tin, the rate being dependent on the severity of the cold-work applied. Tempering after deformation greatly reduces the rate again.—B. Bl.

**\*The Exact Measurement of the Specific Heats of Solid Substances at High Temperatures. VII.—The Calorimetric Behaviour of Zirconium.** F. M. Jaeger and W. A. Veenstra (*Rec. trav. chim.*, 1934, 53, 917-932).—Hysteresis phenomena, already observed in the case of beryllium, are very marked with zirconium. A reversible transformation  $\alpha \rightleftharpoons \beta$  is known to occur in zirconium at 870° C., but further unreproducible complications occur at about 620° C. The temperature varies with the previous treatment and grain-size of the metal. The values for heat content, &c., vary considerably. An attempt is made to explain the complex phenomena observed in the light of Smits' conception of dynamic allotropy, which assumes the existence of two "pseudo components" in movable equilibrium.—C. E. H.

**\*Mechanical Properties of Electrolytic Zinc.** M. D. Zudin (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (7), 116-126; *C. Abs.*, 1935, 29, 4308).—[In Russian.] Soft pure electrolytic zinc assumes, in the process of rolling at

70°–80° C., a fine crystalline structure, which remains unchanged when heated up to 200° C. When it is heated from 200° to 250° C., the crystals grow larger, and a considerable reduction of mechanical properties results. 48 photomicrographs are given.—S. G.

**Effect of Cadmium and of Rapid Cooling on the Structure of Cast Zinc.** M. D. Zudin (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (3), 104–107; *C. Abs.*, 1935, 29, 4305).—[In Russian.] Rapid cooling increased the hardness of pure zinc by 26% and of zinc containing 0.5 and 1% cadmium by 13 and 14%, respectively.—S. G.

**Zirconium and Its Compounds.** Maurice Dérivé (Rev. mat. constr. trav. publ., 1935, 61–63B).—A review.—S. G.

**\*Investigations at Pressures up to 5000 kg./cm.<sup>2</sup>** H. Ebert (*Physikal. Z.*, 1935, 36, 385–392).—Fundamental data relating to the pressure apparatus are discussed, and the following empirical relations for the relative changes of length ( $\Delta l/l$ ), expressed in mm./metre when the samples are subjected to pressures,  $p$  kg./cm.<sup>2</sup>, are deduced for the respective metals at 20° C.; iron (single crystal),  $\Delta l/l = 1.97 \times 10^{-7}p - 0.4 \times 10^{-12}p^2$ ; aluminium (single crystal),  $\Delta l/l = 4.48 \times 10^{-7}p - 0.8 \times 10^{-12}p^2$ ; gold (single crystal),  $\Delta l/l = 1.96 \times 10^{-7}p - 2.0 \times 10^{-12}p^2$ ; lead (single crystal),  $\Delta l/l = 8.00 \times 10^{-7}p - 4 \times 10^{-12}p^2$ ; silver (single crystal),  $\Delta l/l = 3.35 \times 10^{-7}p - 0.5 \times 10^{-12}p^2$ ; magnesium (single crystal),  $\Delta l/l = a \times 10^{-7}p - b \times 10^{-12}p^2$ , the values of  $a$  ranging from 9.68 to 9.77, and of  $b$  from 7.0 to 9.0 according to the inclination of the pressure to the crystallographic axis; copper (polycrystal),  $\Delta l/l = 2.39 \times 10^{-7}p - 0.79 \times 10^{-12}p^2$ ; Manganin (polycrystal),  $\Delta l/l = 2.75 \times 10^{-7}p - 2.0 \times 10^{-12}p^2$ ; steel,  $\Delta l/l = 2.10 \times 10^{-7}p - 1.0 \times 10^{-12}p^2$ . Bridgman's results (where available) are also given. The following values of the cubic compressibilities of the metals ( $K \times 10^7$ ) for  $p = 0$  are deduced: silver, 10.05; aluminium, 13.44; gold, 5.88; copper, 7.17; iron, 5.91; magnesium, 29.09; lead, 24.00; Manganin, 8.25; steel, 6.30.—J. S. G. T.

**Light and Ultraviolet Reflection by Various Materials.** A. H. Taylor (*Trans. Illuminating Eng. Soc.*, 1935, 30, 563–567).—Reflection factors for visible light and ultraviolet radiations at 2967 Å., as measured with the Taylor reflectometer, are tabulated for a number of materials, including aluminium foil and sheet with various finishes, nickel- and silver-plated surfaces, rhodium, and tinned metal.—J. C. C.

**Fifth Report of the Atomic Weights Committee of the International Union for Chemistry.** G. P. Baxter, O. Hönigschmid, P. Lebeau, and R. J. Meyer (*Ber. deut. chem. Ges.*, 1935, [A], 68, (6), 73–85).—The atomic weight of 92.91 is adopted for niobium. Investigations on the atomic weights of carbon, nitrogen, sodium, calcium, krypton, niobium, molybdenum, iodine, and caesium are described, together with Aston's investigations on the rare earth metals and some recent work on tantalum, lead, radium, and protactinium. A complete list of international atomic weights is appended.—P. M. C. R.

**\*A Discussion of the Transition Metals on the Basis of Quantum Mechanics.** N. F. Mott (*Proc. Phys. Soc. (Lond.)*, 1935, 47, 571–588).—The magnetic susceptibilities and electrical conductivities of cobalt, nickel, palladium, and of their alloys with copper, silver, and gold are discussed in accordance with the quantum theory of metals. The number of electrons in the outermost  $s$  state must be 0.6 per atom in the transition metals, and 1 in the noble metals; magnetic properties of the alloys are explained on this basis. A quantum-mechanical explanation of the relatively high resistance of the transition metals is given; the theory is supported by values of the electrical resistances of the alloys.—J. S. G. T.

**\*Direct Effect of Light on the Electrical Resistance of Metals.** A. Etzrodt (*Physikal. Z.*, 1935, 36, 433–441).—Majorana (*Physikal. Z.*, 1932, 33, 947) has claimed to have established the fact that the incidence of light on certain

metals causes the electrical resistance of these metals to be increased, apart from the purely thermal effect produced. E. now shows that the phenomenon observed by M. is purely thermal in character.—J. S. G. T.

**\*A Simple Theory on the Anomaly of Electrical Resistance of Ferromagnetic Substances.** Tokutarō Hirone (*Sci. Rep. Tōhoku Imp. Univ.*, 1935, [1], 24, 122–127).—[In English.] The anomaly is explained by the fact that the mean path of the conduction electron is affected by the rotational vibration of the molecular magnets in the Honda–Okubo model of a ferromagnetic substance. It is to be noted that the equation of corresponding state exists between the reduced resistance and the reduced temperature of the ferromagnetic substance. This conjecture has been confirmed by results of measurements on iron, cobalt, and nickel.—S. G.

**\*Change of the Resistance of Metals in a Magnetic Field at Low Temperature.** W. J. de Haas and P. M. van Alphen (*Comm. K. Onnes Lab. Leiden*, 1933, (225a), 1–15).—[In English.] See *Met. Abs. (J. Inst. Metals)*, 1933, 53, 489.—L. A. O.

**Investigations Relating to the Matteucci Effect.** F. v. Schmoller (*Z. Physik*, 1935, 93, 35–51).—The Matteucci effect—the production of an e.m.f. between the ends of a twisted ferromagnetic wire subjected to a change of magnetization—is investigated in considerable detail. The phenomenon is characterized by two phases of opposite sign corresponding to the direction of change of magnetization; characteristics of these phases are discussed.

—J. S. G. T.

**\*The Direct Measurement of the Peltier Coefficient.** A. J. Woodall (*Proc. Phys. Soc. (Lond.)*, 1935, 47, 615–625).—A new apparatus, including all-metal calorimeters separate from the specimens being tested, and employing a differential platinum resistance thermometer, for the measurement of Peltier coeffs. is described. Employing a sufficiently good thermostatic device, the apparatus is capable of just detecting a steady heating effect of the order of  $10^7$  cal./sec. The Peltier coeff. for copper against Constantan is found to be constant for values of the current between 0.1 and 0.012 amp.—J. S. G. T.

**Thermo-Electric Effect and the Supra-Conducting State.** E. F. Burton, F. G. A. Tarr, and J. O. Wilhelm (*Nature*, 1935, 136, 140).—No thermo-electric effect exists between tin and lead when both metals are in the superconducting state (*i.e.* below  $3.7^\circ$  abs.), but the effect is observed when either one of the junctions is raised above the superconducting point of one of the metals.—E. S. H.

**Asymmetric Rectangular [Hysteresis] Loops Associated with Circular Magnetization.** F. v. Schmoller (*Z. Physik*, 1935, 93, 52–54).—The magnetization curves of a twisted cold-drawn nickel wire subjected to circular magnetization by passage of a direct current show asymmetry about the magnetization axis of the curve which is displaced parallel to the axis of field-strengths. The Barkhausen magnetic effect is also found to be influenced by the torsion.—J. S. G. T.

**\*Magnetic Properties of Metals at Low Temperatures.** W. J. de Haas and P. M. van Alphen (*Comm. K. Onnes Lab. Univ. Leiden*, 1933, (225b), 16–26).—[In English.] See *Met. Abs. (J. Inst. Metals)*, 1933, 53, 489.—L. A. O.

## II.—PROPERTIES OF ALLOYS

(Continued from pp. 370–377.)

**\*Strength Tests of Thin-Walled Duralumin Cylinders of Elliptic Section.** Eugene E. Lundquist and Walter F. Burke (*Nat. Adv. Cttee. Aeronautics, Tech. Notes No. 527*, 1935, 23 pp.).—The shearing stress at failure in torsion for thin-walled elliptical cylinders is equal to that for circumscribed circular

cylinders of the same wall-thickness and length. Since buckling of the walls occurs at the ends of the mirror axis prior to failure, the calculated shearing stress for the elliptical cylinder is analogous to the modulus of rupture. For pure bending in the place of the major axis the calculated stress on the extreme fibre at failure is greater than the corresponding stress for similar circumscribed circular cylinders. Since small imperfections produce wide scattering of the test results the bending strength determinations are somewhat indefinite and the probable value must be estimated after consideration of all the data for both types of cylinders before choosing a design. For combined transverse shear and bending in the plane of the major axis the strength is dependent on the value of  $M/r^2V$  (representing the loading and stress conditions). At low values of  $M/r^2V$  failure occurs in shear and as the value approaches zero (a condition of pure transverse shear) the shearing stress on the neutral axis at failure is about 1.25 times the shearing stress at failure in torsion. At large values of  $M/r^2V$  failure occurs in bending and the strength developed is equal to the lowest strengths developed for similar cylinders in pure bending. At intermediate values of  $M/r^2V$  there is a transition from shear to bending failure. The experimental results are shown in numerous tables and graphs, and a short bibliography is appended.—A. R. P.

\***An Electrochemical Investigation of Solid Cadmium-Antimony Alloys.** (Ölander.) See p. 435.

\***The System Iron-Cobalt-Tin.** Werner Köster and Werner Geller (*Arch. Eisenhüttenwesen*, 1934-1935, 8, 557-560).—The system has been examined by thermal and micrographic methods up to 40% tin. The two compounds  $Fe_2Sn$  and  $Co_2Sn$  form a continuous series of solid solutions. Addition of cobalt increases the temperature of the equilibrium  $\alpha + \text{liquid} = Fe_2Sn$  to meet the equilibria  $\alpha + \text{liquid} = \gamma$  of the cobalt-iron system and  $\text{liquid} = \gamma + Co_2Sn$  of the cobalt-tin system in the 4-phase equilibrium  $\text{liquid} + \gamma = \alpha + Co_2Sn-Fe_2Sn$  solid solution. The solubility of the compounds in the  $\alpha$  and  $\gamma$  phases decreases with decrease in temperature, hence certain alloys are subject to precipitation-hardening, the reheating temperature being 700° C. for  $\alpha$ , 600° for  $\alpha + \gamma$ , and 750° C. for  $\gamma$ -alloys. The magnetic properties, hardness, and microstructure of certain of the alloys are discussed.—A. R. P.

\***The System Iron-Cobalt-Titanium.** Werner Köster (*Arch. Eisenhüttenwesen*, 1934-1935, 8, 471-472).—The system has been studied by thermal and micrographic methods up to 22% titanium; only the two binary compounds  $Fe_3Ti$  and  $Co_3Ti$  exist, and these form a continuous series of solid solutions, the solubility of which in the  $\alpha$ - and  $\gamma$ -phases increases with increase in temperature from about 2% at room temperature so that alloys containing these can be precipitation-hardened by suitable heat-treatment. Thus, after quenching from 1100° C., the alloy with 55% cobalt and 7.6% titanium has a Brinell hardness of 420 which is increased to 600 by tempering at 600° C.; the corresponding values for the alloy with 75% cobalt and 4.5% titanium are 205 and 305. The eutectic equilibrium of the iron-titanium system and the peritectic equilibrium of the iron-cobalt system are displaced to lower temperatures by the addition of the third metal and terminate at 1150° C. in the 4-phase reaction  $\text{liquid} + \alpha\text{-solid solution} = \gamma\text{ solid solution} + Fe_3Ti-Co_3Ti$  solid solution. From this point the equilibrium  $\text{liquid} = \alpha + Fe_3Ti-Co_3Ti$  extends to the eutectic equilibrium of the Co-Ti system, and the equilibrium  $\alpha = \gamma + Co_3Ti-Fe_3Ti$  to room temperature.

—A. R. P.

\* **$\alpha$ -Aluminium-Bronze and Its Mechanical Properties.** P. I. Gradusov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (7), 126-137; *C. Abs.*, 1935, 29, 4315).—[In Russian.]  $\alpha$ -Aluminium-"bronze," containing  $6.5 \pm 0.5\%$  aluminium and the remainder copper, is a good substitute for bronzes con-

taining (1) tin 4, zinc 3, copper 93%, or (2) tin 6.5, phosphorus 0.4, copper 93.1%. The  $\alpha$ -aluminium—"bronze" was melted in an oil-fired furnace. The alloy contained 6.62% aluminium and was used in a series of mechanical tests. Its mechanical strength at temperatures up to 500° C. was 1.5 times that of phosphor-bronze. The  $\alpha$ -aluminium—"bronze" was free from defects. It was rolled, first hot and then cold, into plate, rod, and wire. It has good plastic qualities and is suitable for the production of springs. It is not inferior to phosphor-bronze in corrosion-resistance to air or in solutions, and is also resistant to abrasion. It is recommended as a substitute for ordinary bronze in machine construction, but not in the form of fine wire (e.g. in screens) of 0.30–0.07 mm. diameter.—S. G.

\***Heusler Alloys.** S. Valentiner and G. Becker (*Z. Physik*, 1935, 93, 629–633).—Polemic with Heusler (*Met. Abs.*, 1934, 1, 579). New measurements of the magnetic and electric properties of the alloy containing manganese 25, aluminium 25, and copper 50% confirm the stability of the crystal  $MnAlCu_2$  at about 300° C. and support the authors' contention (*Met. Abs. (J. Inst. Metals)*, 1933, 53, 620) that the magnetic properties of Heusler alloys are associated with the presence of crystals of  $MnAlCu_2$ .—J. S. G. T.

\***Standard Manganin Containing Aluminium, and Its Properties.** P. I. Gradusov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (6), 101–105; *C. Abs.*, 1935, 29, 4315).—[In Russian.] Manganin (copper 85, manganese 12, nickel 3%), in which aluminium replaces nickel, behaves during melting, pouring, and heat-treatment like the ordinary alloy. Its mechanical properties are indistinguishable from those of the standard alloy when its tempering temperature is not above 700° C. The specific resistance of the aluminium-Manganin increases with increase of tempering temperature. In practice it can be used up to 400° C.—S. G.

\***Mechanical Properties of Double Manganin.** P. I. Gradusov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (6), 105–111; *C. Abs.*, 1935, 29, 4315).—[In Russian.] Samples of double Manganin containing copper 71.49, manganese 20.79, and aluminium 6.11% were tempered at 550°, 600°, 650°, 700°, 750° and 800° C., and their mechanical properties were then studied. Heating to 800° C. followed by slow cooling gives the best results.—S. G.

\***The Recovery of the Electrical Resistance of Binary Solid Solution Alloys of Copper, Silver, and Gold from the Consequences of Cold-Work.** G. Tammann and K. L. Dreyer (*Ann. Physik*, 1933, [v], 16, 657–666).—*Cf. Met. Abs. (J. Inst. Metals)*, 1933, 53, 177. The increase in the electrical resistance produced by cold-work and the subsequent reduction by annealing have been measured for solid solution alloys of copper with zinc, tin, nickel, manganese, and aluminium, of silver with zinc, cadmium, and gold, and of gold with silver and copper. The increase is greater the greater the difference between the lattices of the two alloying metals and the greater the atomic radius of the solute metal.—v. G.

\***The Elastic Properties of Gold-Silver Single Crystals.** H. Roehl (*Ann. Physik*, 1933, [v], 16, 887–906).—The elastic constants in  $10^{-13}$  cm.<sup>2</sup>/dyne and the anisotropy of single crystals of gold and silver alloys are :

Atomic-% Gold.	$\Delta_{11}$ .	$\Delta_{44}$ .	$\Delta_{12}$ .	$E_{111}/E_{100}$ .	$G_{100}/G_{111}$ .
0	23.2	22.9	— 9.93	2.72	2.26
25	20.7	20.5	— 8.91	2.66	2.26
50	19.7	19.7	— 8.52	2.65	2.25
75	20.5	20.6	— 9.09	2.68	2.24
100	22.7	22.9	— 10.35	2.72	2.26

**Nickel in the Precious Metal Field.** Eric H. Swanson (*Metals and Alloys*, 1935, 6, 194).—Letter to the Editor. The statement that addition of about 2% of nickel to precious metal alloys containing less than 75% (gold + platinum) seriously impairs the tarnish-resistance is incorrect. Alloys containing about 45% (gold + platinum + palladium) and 4.25% nickel have a high resistance to tarnishing and give excellent castings.—A. R. P.

**Nickel in Dental Alloys.** Wilmer Souder (*Metals and Alloys*, 1935, 6, 194).—Letter to the Editor. Small quantities of nickel in casting alloys for dental restorations appear to reduce their elasticity and resilience; the nickel has a tendency also to produce irritation of the tissues. At least 65% of precious metals is the minimum necessary to obtain tarnish-resistance. (Cf. preceding abstract.)—A. R. P.

†**Beryllium-Copper Alloy.** F. G. Benford (*Gen. Elect. Rev.*, 1935, 38, 297-299).—The heat-treated 2.5% beryllium-copper alloy, whilst possessing excellent mechanical properties has two disadvantages: it is expensive and has a low electrical conductivity (25% that of copper). The addition of a small quantity of cobalt reduces the amount of beryllium necessary for hardening by precipitation. The addition of cobalt not only reduces the cost of the alloy, but also increases the electrical conductivity to about 50% that of pure copper. With the ternary alloys the precipitation-hardening temperature is increased from 275° to about 500° C. Certain compositions have been standardized and the cost of one of these is about  $\frac{1}{3}$  that of the binary beryllium-copper alloy. It is available in all forms and has been applied to a variety of uses. [Note by abstractor: No actual compositions are given.]—S. V. W.

\***Effect of Low Temperatures on the Structure and Mechanical Properties of Antifriction Alloys.** S. S. Nekruitui and B. D. Grozin (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (3), 91-104; *C. Abs.*, 1935, 29, 4314).—[In Russian.] The behaviour at low temperatures of antifriction alloys used in railway cars and locomotives was studied. The alloys tested were Sharpe Babbitts B-1, B-2, B-3, B-4, and B-5 and a lead-base calcium-Babbitt containing calcium 0.75-1.1, sodium 0.7-1.0, antimony less than 1%, and other elements. Samples, first cooled to -53° or -75° C., then heated in oil or air to +12°, +70°, or +75° C., were tested for hardness, impact, compression, and abrasion, and were also examined microscopically. Low temperatures down to -50° C. have no harmful effect on the tin or calcium-Babbitts, but below this temperature a softening effect is produced, owing to the decomposition of Sn-Sb and Pb<sub>2</sub>Ca crystals in the Sharpe or calcium-Babbitts, respectively. Calcium-Babbitt compares well with the Sharpe standard Babbitt B-1.

—S. G.

**Notes on Heat-Resisting Metals.** N. P. Inglis (*J. Soc. Glass Tech.*, 1933, 17, 366-377).—Deals with developments and improvements in various metals to improve their heat-resistance. The extreme susceptibility of the Nichrome type alloys to gases containing sulphur is shown, and the effect of an addition of aluminium to these alloys is dealt with. I. quotes experimental results showing the great increase in resistance to sulphur-containing gases which is obtained by an addition of aluminium to the Nichrome alloys.—S. G.

\***The Optical Constants of the Copper-Nickel Alloys.** H. Lowery, J. Bor, and H. Wilkinson (*Phil. Mag.*, 1935, [vii], 20, 390-409).—The optical const. of 12 copper-nickel alloys containing, respectively, 0, 8.8, 18.4, 24.7, 35.2, 45.1, 53.2, 64.9, 76.0, 85.5, 91.8, and 100% (atomic) of nickel were measured, using a polarimetric method, for wave-lengths 5780, 5461, and 5348 Å. Stress is laid on the method of polishing adopted, the object of which is the production of a very thin flowed layer. The significance of the results is briefly discussed.

—J. S. G. T.



“K” Monel Metal. — (*Mech. World*, 1935, 97, 632).—By the addition of aluminium and careful adjustment of composition, “K” Monel metal is obtained, the properties of which may be widely varied by suitable heat-treatment. Examples of the variation in properties obtainable by specified heat-treatment, &c., are given.—F. J.

\*Magnetic Investigations Relating to Precipitable Iron-Nickel Alloys. F. Preisach (*Z. Physik*, 1935, 93, 245-268).—The changes of magnetic properties of 2 iron-nickel alloys of respective compositions nickel 55, iron 45% with 1% of beryllium added to the melt, and nickel 38, iron 46, copper 15, manganese 1% (Isoperm) during the process of precipitation-hardening, are investigated. The results are interpreted on the assumption that in addition to the amount and distribution of the precipitated material, the recovery of the crystal lattice is of extreme importance in the discussion of the development of magnetic hardening. Observations referring to the inter-relationship of the precipitation process and crystal lattice recovery are discussed in considerable detail.

—J. S. G. T.

Wear of Iron[-Nickel] Alloys on Emery Paper and Their Hardness. Willi Tonn (*Arch. Eisenhüttenwesen*, 1934-1935, 8, 467-470).—The comparative rates of wear of metals or alloys may be determined by holding specimens under a load of 1 kg. on the surface of a rotating disc covered with emery paper and moving the specimen slowly along a diameter so that it continuously meets fresh emery. The loss in weight is roughly proportional to the scratch hardness, and is not affected by heat or mechanical treatment. The wear of iron-nickel alloys decreases rapidly with increasing nickel content to 30%, then remains constant to 70% nickel after which it increases slightly; the Brinell hardness curve shows a well-marked maximum at 17% nickel, and is practically a horizontal line between 35 and 100% nickel.—A. R. P.

\*The Causes of the Change in Length of Invar Wires. F. Neumann and H. Johannsen (*Z. Instrumentenkunde*, 1934, 54, 173-190).—The behaviour of carefully treated and well-aged Invar Jäderin wires depends on (1) changes in structure which lead to slight expansion, and (2) accidental strains in the wires. The paper deals principally with the latter, and describes investigations on the effects of artificially produced kinks on the dimensions of the wire. The dimensional changes are related to the geometrical form of the kinks. In a wire under tension there is an initial shortening when a kink is formed, but this is followed by a lengthening. Recommendations are made for the handling and usage of Invar wires to give the most accurate results from experiments in which they are used.—C. E. H.

\*The System Nickel-Manganese. Siegfried Valentiner and Gotthold Becker (*Z. Physik*, 1935, 93, 795-803).—Nickel-manganese alloys, quenched from a high temperature, are found to consist of a solid solution of manganese in nickel, or, in the case of alloys containing over 80% manganese, of a solid solution of nickel in  $\beta$ -manganese. A tetragonal phase is present in annealed alloys containing from 40 to 70% manganese. The electrical conductivities of alloys containing, respectively, 25 and 50% manganese increase considerably when the alloys are annealed at about 450° C.; the temperature coeff. of electrical conductivity of these alloys increase under the same condition. The effect is attributed to the replacement of  $Ni_3Mn$  in the quenched alloys by  $NiMn$  in the annealed alloys. The effect of the two component metals is clearly brought into evidence in the magnetic properties of the alloys. Alloys containing about 25% manganese are very strongly paramagnetic; this phenomenon is attributable to the presence of the superstructural alloy  $Ni_3Mn$ . Ferromagnetism is not exhibited by the superstructural alloy containing about 50% manganese; this is attributable to the presence of a tetragonal crystal lattice. Further, these latter alloys are

characterized by increased values of electrical conductivity. The presence in them of NiMn is confirmed by X-ray analysis.—J. S. G. T.

†**Alloys with a Zinc Base.** Willi Claus (*Z.V.d.I.*, 1935, 79, 385–386).—Advances made in recent years in zinc-base alloys have resulted in greatly increasing their sphere of usefulness in engineering technique. The tensile properties of zinc-base die-castings and the effect of impurities thereon are discussed.—K. S.

\***The Mechanical Properties of Metals at Low Temperatures.** W. Schwinning (*Z.V.d.I.*, 1935, 79, 35–40).—The notched-bar impact strength and endurance properties of numerous metals and alloys at low temperatures are tabulated, and the effects of temperature changes are shown graphically. The tensile properties and endurance bending strength of light alloys, especially of hardened Duralumin and Aldrey, are affected only slightly by cooling to  $-40^{\circ}\text{C}$ . The tensile strength and yield-point of copper and steel are considerably increased and the endurance bending strength is raised by double as much as the tensile strength on cooling to  $-40^{\circ}\text{C}$ .—K. S.

†**Brittleness in Ductile Engineering Materials.** W. E. Lewis (*Trans. Inst. Eng. Ship. Scotland*, 1934–1935, 78, 444–484).—L. deals with the embrittlement occurring in various ductile materials, including the "season-cracking" of cold-worked brass, failure of mild steel in contact with brazing solder, embrittlement occurring when molten tin or tin-lead solders are applied to copper or brass when stressed in tension, and electrochemical disintegration in brass or manganese-brass propellers. In the discussion considerable space is devoted to the Izod impact test as a means of determining brittleness in materials.—J. W. D.

\***The Norbury Relation in the Thermal Resistance of Metals.** H. Reddemann (*Ann. Physik*, 1934, [v], 20, 502–512).—The increase in electrical resistance of alloys is proportional to the increase in thermal resistance up to a high content of alloying metal. This rule has been confirmed by data taken from the literature on alloys of copper, aluminium, gold, silver, platinum, and palladium. The thermal resistance of alloys can, therefore, be calculated in most cases from their electrical resistance.—v. G.

\***Supra-Conducting Alloys.** K. Mendelssohn and Judith R. Moore (*Nature*, 1935, 135, 826–827).—Differences in the behaviour of superconducting alloys in comparison with pure metals have been examined in the case of a lead-bismuth alloy. As an explanation, it is suggested that the temperature at which superconductivity is observed is high in some parts of the alloy while the main part has the same value as pure metals.—E. S. H.

\***Magnetic Properties and Critical Currents of Supraconducting Alloys.** J. N. Rjabinin and L. W. Shubnikow (*Nature*, 1935, 135, 581–582).—The relation between the intensity of the field applied and the magnetic induction produced in alloys of lead-thallium ( $\text{PbTl}_2$ ) and lead-bismuth (lead 65%) at very low temperatures is described. Above a certain critical field-strength these alloys lose their superconductivity.—E. S. H.

\***On the Theory of Transformation of Metallic Solid Solutions.** G. Borelius (*Ann. Physik*, 1934, [v], 20, 57–74).—Equations have been developed on thermodynamic principles for expressing the course of transformations which take place over a temperature range and are subject to hysteresis. The results are applied to the transformation of AuCu and  $\text{AuCu}_3$ .—v. G.

\***Internal Equilibria in Solid Phases.** A. Smits (*Physikal. Z.*, 1935, 36, 367–371).—S. directs attention to the priority of his work on equilibria and heterogeneous transformations in solid phases (*Physikal. Z.*, 1930, 31, 376; and *Met. Abs.*, this vol., p. 155), and discusses subsequent work by Juste and von Laue (*Met. Abs.*, this vol., p. 155) and by Eucken (*Met. Abs.*, this vol., p. 219).—J. S. G. T.

## III.—STRUCTURE

(Metallography; Macrography; Crystal Structure.)

(Continued from pp. 378–380.)

†Questionnaire on Dendrites, the Monocrystalline State, Metal Grains, and the Mechanism of Grain-Growth.—I.—II. Léon Dlougatch (*Rev. Mét.*, 1935, 32, 23–31, 85–99).—(I—) The views of 24 authorities are given and are followed by a brief *résumé* of the replies received on the subject of dendrites (see also abstract from Russian, *Met. Abs.*, 1934, 1, 578). (II.—) The replies received on the subject of the monocrystalline state are reviewed. On the subject of twinned crystals there is some divergence of opinion. While many of the authorities regard twinned crystals as something different from monocrystalline, several point out that it is a matter which may be considered from various aspects and with different results. The various views on grain in alloys and on grain-growth are also reviewed.—H. S.

**The Kinetics of Crystallization Processes.** A. Huber (*Z. Physik*, 1935, 93, 227–231).—A mathematical theory of crystallization is presented, which is a development of that due to v. Göler and Sachs (*J. Inst. Metals*, 1932, 50, 671) and which enables the theoretical conclusions of the latter to be compared with experimental results.—J. S. G. T.

**\*Crystal Growth and the Production of Crystal Nuclei.** I. N. Stranski and R. Kaischew (*Physikal. Z.*, 1935, 36, 393–403).—A mathematical theory of cubical crystal growth based on considerations of energy is developed.

—J. S. G. T.

**\*The Orientation of Crystals in Metals and [its Effect on] Solubility.** Vincenzo Caglioti (*Atti IV Congr. naz. chim. pura applicata* (1932), 1933, 442–446; *C. Abs.*, 1935, 29, 4239).—Out of 99.95% aluminium, repeatedly forged and reheated, 2 small cubes were cut in a direction parallel to that of forging, and were then rolled down to 99%, one being rolled in all directions and the other in one direction only. X-ray examination showed random orientation of the crystals in the first, while, in the second, one (335) axis was substantially parallel and one (135) axis substantially normal to the direction of rolling. The two sheets, 10 cm.<sup>2</sup> in area, were polished and then cleaned by dipping for 10 minutes in 4-N hydrochloric acid. They were then immersed each in 3 litres of 2.5-N hydrochloric acid at 19.8° C., and the hydrogen evolved was measured every 15 minutes up to 4 hrs. The second sheet (uniform orientation of crystals) showed a longer period of induction and a slightly lower rate of solution than the first (random orientation). This may be owing either to a difference of potential of solution for the various faces, or to a regular distribution of an impurity in some of the crystal faces.—S. G.

**\*X-Ray Tests of the Recrystallization of Duralumin.** N. K. Kozhina (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1934, (17), 3–16).—[In Russian, with German summary.] See *Met. Abs.*, this vol., p. 221.—S. G.

**\*X-Ray Determination of the [Crystal] Structure of FeAl<sub>3</sub>.** E. Bachmetew (*Z. Krist.*, 1934, 89, 575–586).—Cf. *Met. Abs.*, this vol., p. 99. The alloy FeAl<sub>3</sub> is shown to possess rhombic crystal symmetry. The crystal lattice constants have the respective values:  $a = 47.43 \text{ \AA}$ ,  $b = 15.46 \text{ \AA}$ ,  $c = 8.08 \text{ \AA}$ . The number of atoms per cell ( $N_A$ ) is 400; the calculated density is 3.811 [temperature not specified]. The crystals belong to the translation group  $\Gamma_6$  (general face-centred) and to space group  $V_6^{23}$ .—J. S. G. T.

**Native Copper.** Antoni Laszkiewicz (*Arch. Mineral. Tow. Nauk. Warszaw.*, 1933, 9, 1–17).—Crystallographic.—S. G.

**\*Growing Magnesium Crystals.** M. Straumanis (*Z. Krist.*, 1934, **89**, 487-493).—Cf. *Met. Abs.*, this vol., p. 15. The growth of magnesium crystals deposited from the vapour phase in hydrogen at a pressure of 0.001-360 mm. of mercury is described and illustrated. During such deposition, in addition to crystals having the equilibrium form bounded by the planes 0001, 10 $\bar{1}$ 0, and 10 $\bar{1}$ 1, others bounded by some of these planes and a curved surface, having a bright and molten appearance are found to occur. Microscopic investigations show that during the initial process of crystal growth, more especially when the vapour is highly supersaturated, crystal nuclei having no external planes can be produced. With increasing size of the nuclei the equilibrium plane surfaces begin to appear, with so-called "Vizinal" surfaces appearing during the intermediate stage. These last surfaces occur only in rapidly growing crystals. As conditions of growth approach equilibrium conditions the "Vizinal" surfaces disappear. In addition to these surfaces and the plane 10 $\bar{1}$ 3 which appears sporadically, no other surfaces were found to characterize non-equilibrium conditions of growth.—J. S. G. T.

**\*Gold-Copper Alloys, more Especially at High Temperatures.** L. Vegard and Arne Kloster (*Z. Krist.*, 1934, **89**, 560-573).—The crystal structures of gold-copper alloys containing respectively the pure components and 75, 50, and 25% (atomic) of the components were determined by X-ray analysis. At 475° C. the alloys are characterized by random atomic distribution; in accordance with previous work, compounds such as CuAu and CuAu<sub>3</sub> cannot be present at this temperature. Although the system copper-gold at 475° C. forms a complete series of true solid solutions (Mischkristalle) the additive law of lattice dimensions is not very closely obeyed. At room temperature the system of alloys shows no lines associated with the presence of metallic compounds, but exhibits the character of solid solutions with random atomic distribution; alloys containing about 50% (atomic) of the constituent metals are characterized by such a conspicuous absence of micro-structure that the X-ray lines, even at large dispersion, almost disappear altogether. Lattice dimensions, so far as they could be determined at room temperature, exhibit the same kinds of departure, as regards magnitude and direction, from the additive law, as at high temperature.—J. S. G. T.

**\*X-Ray Investigation of Palladium-Gold Alloys Saturated with Hydrogen.** H. Mundt (*Ann. Physik*, 1934, [v], **19**, 721-732).—X-ray examination of palladium-gold alloys saturated with hydrogen showed that they are similar to the corresponding silver alloys (see *Met. Abs. (J. Inst. Metals)*, 1933, **53**, 190).—v. G.

**\*Lattice Parameters of Solid Solutions in Silver.** William Hume-Rothery (*Nature*, 1935, **135**, 1038).—The addition of cadmium, indium, tin, and antimony to silver changes its lattice structure to an extent which is proportional to the valency of the added element.—E. S. H.

**\*The Twinning of Single Crystals of Tin.** Bruce Chalmers (*Proc. Phys. Soc. (Lond.)*, 1935, **47**, 733-746).—Methods for preparing single crystals of tin are described, and the crystallographic orientation of the crystals determined optically, the method employed being based on the character of the etched crystal surface. Conditions in which parts of the crystal can be caused to twin either by impact or tension are investigated, and the determination of the energy relations for controlled cases of twinning is described. When twinning occurs, energy equal to  $8 \times 10^5$  ergs is transformed into heat per cm.<sup>3</sup> of crystal twinned. The mechanism of twinning in tin is discussed. The twinning plane is of the 301 type, of which there are 4 sets in the crystal lattice. A sufficient component of sheer stress in the right direction in a set of 301 planes is essential before twinning can occur. When tension is applied in a direction nearly parallel to the 001 axis the result may be either twinning or glide. When there is a component of shear stress in the appropriate

direction in two or more of the twinning planes, twinning occurs on the plane in which the component stress is greatest.—J. S. G. T.

**\*Investigation of Slip under Shear in Single Crystals of Tin.** K. Bausch (*Z. Physik*, 1935, 93, 479–506).—Single crystals of tin were so mounted by soldering in a Polanyi extensometer that thrusts in the slip-planes and slip directions were operative. In the slip-planes (110) and (100) slip occurred in the direction [001] whatever the direction of application of the tensile stress in the planes. The initial parts of the hardening curves followed the cosine law of thrust stress; in subsequent parts intensified hardening was present. In the plane (110) slip also occurred in the direction  $[\bar{1}11]$ ; in the plane (100) slip also occurred in the direction [010]. The mechanism of plastic deformation of crystals is discussed in considerable detail.—J. S. G. T.

**\*A Special Type of Unstable Solid Solutions With Anomalous Lattice Constants.** G. Natta (*Naturwiss.*, 1935, 23, 527–528).—Very thin films of platinum alloys are obtained by immersing silver or copper plates in a very dilute solution of platinum tetrachloride and then treating the deposit with nitric acid. Films  $10^{-6}$  cm. thick afford good electron diffraction patterns which show that they consist of alloys the nature of which depends on whether the ratio of the atomic radii of the deposited and depositing metals is or is not smaller than 1. When platinum, palladium, or silver are deposited on copper the diffraction patterns are those of a cubic face-centred solid solution. When platinum is deposited on silver, however, the freshly-formed film has the same lattice parameter as silver, but this becomes smaller on keeping at room temperature for some days or on heating at  $700^{\circ}\text{C}$ . for a few seconds, indicating that unstable silver solid solutions with up to 50% platinum can exist having the same lattice parameter as pure silver. The change from the unstable loose lattice into the stable compact lattice is often accompanied by a change in colour.—B. Bl.

**\*Statistical Theory of Superlattices.** H. A. Bethe (*Proc. Roy. Soc.*, 1935, [A], 150, 552–575).—The degree of order in an alloy crystal containing two sorts of atoms in equal proportion is calculated statistically, assuming interaction between nearest neighbouring atoms only. At high temperatures the state of the crystal as regards order is similar to that of a liquid. At low temperatures the crystal as a whole is ordered; the structure is “solid-like.” The order at long distances is restricted to two or more dimensions. The long-distance order vanishes at a critical temperature,  $\theta$ . All the physical quantities, plotted as functions of  $T$ , exhibit a kink at  $T = \theta$ , but no jump. This is owing to the fact that at  $\theta$ , two “symmetrical” states exist, having the same energy. The derivatives of physical quantities, e.g. specific heat, exhibit jumps at the critical temperature. The extra specific heat due to the destruction of order is rather large on the low-temperature side of  $\theta$ ; it is 70% of the ordinary specific heat due to thermal motion of the atoms (provided all atoms of the crystal take part in the transition. On the high-temperature side of  $\theta$  it decreases to about 10% of this value.—J. S. G. T.

**\*Intensity Measurements on Deformed Crystals and Solid Solutions.** J. Hengstenberg (with H. Mark, and — Wassermann) (*Ergebnisse tech. Röntgenkunde*, 1931, 2, 139–150; *Brit. Chem. Abs.*, 1934, [A], 1161).—The change in intensity of X-ray reflection for deformed crystals of tantalum, tungsten, molybdenum, or potassium chloride is proportional to the degree of deformation. In gold-silver solid solutions (4, 8, and 10 wt.-% gold) an increase in the diffuse scattered radiation was observed, the Laue intensity distribution being confirmed. Duralumin showed characteristic effects for both solid solution and deformed crystals.—S. G.

**\*Laue Diagrams of Deformed Crystals.** W. F. Berg (*Z. Krist.*, 1934, 89, 587).—[In English.] The following facts relating to Laue X-ray diagrams of deformed crystals are explained: (1) Laue spots of a deformed crystal

are radially elongated if the Laue diagram is taken by transmitted X-rays, (2) the spots are smeared out in all directions in a diagram taken by reflected X-rays, (3) the Laue spots on a diagram of a crystal deformed by bending round an axis normal to the incident beam of X-rays, are elongated in a direction perpendicular to the axis of deformation, (4) a Laue diagram taken by reflected X-rays is comparatively insensitive to errors in setting of the crystal.—J. S. G. T.

**The Laws of Valence-Electron Concentration in Binary Metallic Alloys [Hume-Rothery Rule].** Harald Perlitz (*Acta et Comm. Univ. Tartuensis*, 1933, [A], 24, (2), 16 pp.; *C. Abs.*, 1935, 29, 4719).—Cf. *Met. Abs. (J. Inst. Metals)*, 1933, 53, 629 and *Met. Abs.*, 1934, 1, 297. The rule proposed by Hume-Rothery and by Westgren that the valence-electron concentration, or ratio of the number of valence electrons to the number of atoms, is a definite constant characteristic of each lattice type, is shown to apply to  $\beta$ -,  $\gamma$ -, and  $\epsilon$ -alloys. The values of the constants are 3 : 2, 21 : 13, and 7 : 4, respectively. Since the constant is always greater than 1, a necessary condition for the existence of these structures is that one of the metals contributes at least 2 valence electrons and the other component contributes not more than 1 valence electron to the lattice structure. From the actual existence and non-existence of the alloys that can possibly be formed on the basis of this requirement, it is further concluded that one of the components must be a transition metal with the electron effect, or else either copper, silver, or gold; and that the other component must be a metal of the B-subgroup, or else either beryllium or magnesium. If it be assumed that the intermediary phase with the  $\beta$ -,  $\gamma$ -, or  $\epsilon$ -structure has a stoichiometric formula, and further, on the basis of its magnetic susceptibility, that the valence-electron concentration of silver in its alloys is one, then the valence-electron concentration of the metals having the electron effects, *viz.* manganese, iron, cobalt, nickel, rhodium, palladium, and platinum, is zero, and that of copper, silver, gold, beryllium, magnesium, zinc, cadmium, aluminium, germanium, tin, and antimony is equal to the salt valence, while that of mercury and indium is 4.—S. G.

**Identification of Crystalline Substances by Means of X-Rays.** B. E. Warren (*J. Amer. Ceram. Soc.*, 1934, 17, 73-77).—Methods for the preparation of X-ray diffraction patterns of crystalline materials, the significance of an X-ray diffraction pattern, and its relation to the sample material are discussed. Methods are also given for identification by (a) direct analysis of pattern, and (b) comparison with diffraction patterns of known materials. Difficulties in classifying and cataloguing diffraction patterns are dealt with.—S. G.

**\*A Simple Determination of the Space-Orientation of Crystals.** H. Ekstein and W. Fahrenhorst (*Z. Krist.*, 1934, 89, 525-528).—A graphical method is described for determining crystal orientation from results of X-ray analysis by the rotating crystal method and an optional Laue diagram. The case of a cylindrical crystal is more particularly discussed, but the method is applicable generally. Results relating to cadmium crystals are illustrated.

—J. S. G. T.

**\*A Graphical Method for Indexing [X-Ray] Spectra Taken by the Rotating Crystal Method.** N. J. Seljakow and E. I. Sovz (*Z. Krist.*, 1934, 89, 601-606).—See also *Met. Abs.*, this vol., p. 101. A graphic method is discussed for indexing lines in X-ray spectra taken by the rotating crystal method, rotation taking place about an arbitrary axis. The method is illustrated by reference to lines in the spectrum of a single-crystal of aluminium.

—J. S. G. T.

## IV.—CORROSION

(Continued from pp. 381-384.)

**\*Structural Corrosion of Aluminium Alloys. I.—Electrode Potentials. II.—Structural Corrosion.** G. V. Akimov and A. S. Oleschko (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1934, (18), 3-30).—[In Russian, with English summary.] See *Met. Abs.*, 1934, 1, 582, and this vol., p. 58.—S. G.

**Behaviour of Copper on Exposure to the Elements.** — (*Roy. Inst. Brit. Architects Advance Copy*, 1934, Apr., 4 pp.).—Brief notes are given on the formation of natural and artificial patina, on heat bronzing, and on the action of various natural waters, on copper.—A. R. P.

**\*Corrosion of Magnesium Alloys. II.—Corrosion of Cast Magnesium Alloys.** W. O. Kroenig and G. A. Kostylew (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1933, (4), 5-58; *C. Abs.*, 1935, 29, 4724).—[In Russian, with English summary.] Cf. *Met. Abs. (J. Inst. Metals)*, 1933, 53, 195. Samples of cast magnesium containing up to aluminium 12.21, manganese 1.16, zinc 3.99, copper 2, cadmium 4.95, nickel 3.10, silicon 0.69, and lead 0.82% were tested for corrosion in solutions of chloride salts (including sea-water), in distilled and tap-water and in air in atmospheric conditions. The method of measuring corrosion was the same as that used in the previous work. In sea-water magnesium containing manganese is more resistant than when pure, the optimum amount of manganese being 0.3-0.4%; in distilled and tap-water manganese has no effect. Addition of aluminium increases the resistance of magnesium to sea-water and particularly to distilled and tap-water; the optimum aluminium content in the latter case being 6-8%. Addition of manganese to magnesium-aluminium alloys increases their resistance to sea-water, the optimum manganese content being 0.3-0.4% for all magnesium-aluminium alloys; in distilled and tap-water manganese has no effect on these alloys. Cadmium has a beneficial effect on binary magnesium alloys in distilled water, when its content is 5%, but no effect on ternary alloys. Zinc has no effect on the corrosion of magnesium alloys, but its use is desirable because it improves the mechanical properties of ternary magnesium alloys. Lead and nickel are definitely harmful. Copper in amounts greater than 1% is harmful to binary alloys, but when only 0.1% of copper is added to magnesium alloys with aluminium, zinc, and manganese it is somewhat beneficial, particularly in the first period of attack. Beryllium (0.25-0.75%) has no advantage over the other additions to magnesium. Calcium has no effect on magnesium or its alloys. Commercial magnesium, as well as its manganese, cadmium, and zinc alloys, when etched in acid solutions containing dichromates did not become more resistant in water or sea-water, but with magnesium-aluminium alloy the resistance increased 600-700 times. It is concluded that none of the magnesium alloys investigated is sufficiently resistant to corrosion by sea-water. In distilled tap water the best composition is aluminium 6-8, zinc 1-1.5, and manganese 0.3-0.4%. Etched cast Elektron is very resistant in air in atmospheric conditions. A bibliography of 49 references is given.—S. G.

**\*Corrosion of Magnesium Alloys. III.—Influence of Heat-Treatment on the Corrosion of Cast Magnesium Alloys.** W. O. Kroenig and G. A. Kostylew (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1933, (5), 5-30; *C. Abs.*, 1935, 29, 4724).—[In Russian, with English summary.] Magnesium alloy castings containing up to aluminium 8.38, manganese 1.16, zinc 3.99, copper 0.51, cadmium 4.95, iron 0.11, nickel 3.10, and silicon 0.69% were heat-treated in an electric muffle furnace. The samples were heated to 300°, 400°, and 500° C. and kept at these temperatures for 24-32 hrs. Some of the samples were then quenched in water and some were annealed for 15 hrs. at 150° C. Corrosion tests on these samples were carried out in a 3% sodium

chloride solution. Heat-treatment decreased corrosion-resistance in nearly all cases. Microscopic analysis of the heat-treated samples revealed that, unlike aluminium alloys, the magnesium alloys become, in the hardening process, less homogeneous in structure because of the formation of small inclusions distributed through the solid solutions of the alloys.—S. G.

**\*Corrosion of Magnesium Alloys. IV.—The Corrosion of Magnesium and Elektron in Contact with Other Metals.** W. O. Kroenig and G. A. Kostylew (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1933, (7), 5–24; *C. Abs.*, 1935, 29, 4724).—[In Russian, with English summary.] Samples of magnesium containing aluminium 0.03, iron 0.04, and silicon 0.02% and of Elektron containing aluminium 8.4, iron 0.06, silicon 0.02, copper 0.12, zinc 1.02, and manganese 0.36% were placed together with samples of other metals in a 3% solution of sodium chloride, the circuit between the two dissimilar metals was then closed and corrosion measured by the amount of hydrogen liberated at the magnesium or Elektron cathode. Similar experiments were carried out in 0.5% solutions of potassium chloride, sodium chloride, magnesium chloride, potassium nitrate, magnesium sulphate, potassium sulphate, and sodium sulphate. In all cases the corrosion of magnesium or Elektron was greater than when placed alone in the solutions. Regarding the promotion of corrosion in 3% sodium chloride solution, the elements, arranged in order of decreasing effect, are: platinum, aluminium, iron, nickel, copper, lead, manganese, zinc, and mercury. Intensity of corrosion was directly proportional to the size of the magnesium or Elektron sample, up to a certain ratio between its size and that of the other metal after which the intensity decreases somewhat; it is also directly proportional to the distance between the electrodes.—S. G.

**\*Corrosion of Wrought Aluminium Alloys.** W. O. Kroenig and N. D. Bobovnikov (*Trans. Research Inst. Aircraft Materials (U.S.S.R.)*, 1934, (16), 4–39; *C. Abs.*, 1935, 29, 4723).—[In Russian, with English summary.] A study was made of the effect of heat-treatment on mechanical properties and resistance to corrosion of (1) alloys of the Duralumin type containing varying amounts of copper, manganese, magnesium, and silicon; (2) magnesium-cadmium-aluminium and magnesium-cadmium-silicon-aluminium alloys, and (3) alloys of the Magnalium type (magnesium up to 8 and manganese up to 2%). Samples were heated to 500° C. in a bath of molten salt and then either quenched in water or aged artificially at 150° C. for 3, 6, 12, or 24 hrs. or naturally at room temperature for 7 days. It is concluded that when the copper content in Duralumin exceeds 5–5.2%, heat-treatment results in a decrease in mechanical properties. Alloys of the Duralumin type containing 0.5–0.7% silicon are improved mechanically by artificial ageing and to a lesser degree by ageing at room temperature. The same alloys acquire a greater resistance to corrosion after natural ageing, but artificial ageing has an opposite effect because of separation of CuAl or MgSi at the grain boundaries. It is possible to regulate the separation of these compounds by changing the ageing temperature and the duration, but this will lead to a decrease in mechanical properties. The addition of cadmium to aluminium-magnesium alloys does not improve their mechanical properties or increase their resistance to corrosion either before or after heat-treatment. Alloys containing up to 8% magnesium and 1% manganese possess high mechanical properties and good resistance to corrosion, either before or after heat-treatment; the best composition, however, is 6% magnesium at 0.5–1.0% manganese.

—S. G.

**\*On Corrosion Phenomena of [Chromium-Nickel] Wire-Wound Radio Resistances on Steatite Insulators.** E. Schürmann and W. Esch (*Elektrotech. Z.*, 1934, 55, 1003–1005); and *Mitt. Material., Sonderheft* 26, 1935, 149–151).—Corrosion phenomena have been frequently observed on the chromium-



nickel alloy wires wound on steatite insulators for radio resistances. The cause of these phenomena has been traced to the presence of pyrite crystals in the steatite; comparative tests on the reaction between powdered pyrite and nickel sheet and the electrochemical nature of the corrosion have confirmed this conclusion.—B. Bl.

**\*The Causes of the Corrosion of the [Nickel and Constantan] Wires in Heating Pads.** M. Popp (*Kautschuk*, 1935, 11, 60–61; *C. Abs.*, 1935, 29, 4725).—Experiments by P. indicate that the explanation advanced by Schürmann and Esch (*Met. Abs.*, 1934, 1, 583) for the corrosion of nickel by asbestos is erroneous. No iron sulphide could be detected, either chemically or microscopically, in any asbestos at the disposal of P., nor does the literature mention this impurity, and oxidation with hydrogen peroxide, nitric acid, and potassium chlorate–hydrochloric acid did not give any evidence of sulphuric acid. The green oxidation products on nickel and Constantan wires in contact with asbestos are chiefly nickel chloride and cupric chloride with smaller proportions of nickel carbonate and copper carbonate. Various commercial grades of asbestos contain chlorine, part of which is soluble in water, and more of which is soluble in aqueous sodium hydroxide. The water exts. also contain magnesium, calcium, sodium, potassium, and silicon and are in some cases alkaline, in others acid, to turmeric. The samples which contain the most soluble magnesium are alkaline. Some of the chlorine of asbestos can be extracted only with acids. Accordingly, asbestos probably contains alkali chlorides, magnesium chloride, and magnesium oxychloride. Magnesium chloride is the agent responsible for the attack on the metal, by virtue of the solution of the hydrochloric acid in the moisture present, and its electrolysis to hydrochloric acid by the electric element formed with the magnetite present as an impurity. (See also following abstract.)—S. G.

**The Causes of the Corrosion of the Wires in Heating Pads.** E. Schürmann and W. Esch (*Kautschuk*, 1935, 11, 78–79; *C. Abs.*, 1935, 29, 4725).—Cf. preceding abstract. Iron sulphide in asbestos was reported by Dammer and Tietze (*Die nutzbaren Mineralien*, 1914, 2, 246, 249). Popp offers no evidence to invalidate the conclusions of S. and E., and the part played by magnesium chloride should be substantiated by experiments with asbestos containing no magnetite.—S. G.

**\*Solution Rates of Zinc Electrodes in Acid Solutions.** H. Mouquin and W. A. Steitz (*Trans. Electrochem. Soc.*, 1935, 67, 339–344; discussion, 344–345).—See *Met. Abs.*, this vol., p. 162.—S. G.

**\*Atmospheric Exposure Tests on Non-Ferrous Screen Wire Cloth.** G. Willard Quick (*J. Res. Nat. Bur. Stand.*, 1935, 14, 775–793; and *Nat. Bur. Stand. Research Paper No. 803*).—Atmospheric corrosion tests on 7 compositions (90 : 10 and 80 : 20 copper–zinc; 75 : 20 : 5 copper–nickel–zinc; 70 : 30 nickel–copper; unalloyed copper; 98 : 2 copper–tin, and 95 : 5 copper–aluminium) of non-ferrous screen wire cloth were carried out by the (U.S.) National Bureau of Standards in co-operation with the American Society for Testing Materials over a period of about 9 years. The specimens were exposed at Pittsburgh, Pa., in a heavy-industrial atmosphere; at Portsmouth, Va., and Cristobal, C.Z., which have temperate and tropical sea-coast atmospheres, respectively, with some industrial contamination, and at Washington, D.C., having a normal inland atmosphere. At the end of the tests at least one failure had occurred in each of the materials exposed at Pittsburgh; one of the materials (80 : 20 copper–zinc) had failed at both Portsmouth and Cristobal; whilst at Washington none of the materials had failed. The tensile strengths of the materials before and after exposure are compared. There was no consistent relation between the results of laboratory accelerated-corrosion tests and the atmospheric exposure tests at any of the locations.—S. G.

**Corrosion in Sea-Water in Which the Action of the Lower Forms of Life Play a Part.** C. Bärenfänger (*Wasser*, 1934, 8, (II), 44-48; *C. Abs.*, 1935, 29, 4496).—The action of the lower sea plants and animals on metal and wood in sea-water is briefly discussed. A lacquer having a benzylcellulose base is reported to be particularly effective protection against both biological and chemical action. Tests show that this lacquer protects iron from the action of sea-water saturated with air and carbon dioxide.—S. G.

**Corrosion by Warm Water and Its Prevention.** L. W. Haase (*Wasser*, 1934, 8, (II), 111-123; *C. Abs.*, 1935, 29, 4496).—The action of warm water (50°-85° C.) on metals and protective coatings used for pipes, various types of warm-water plants, and methods of treatment to prevent corrosion are discussed.—S. G.

**\*Motor-Electrolytic Current as a Factor in Corrosion.** E. Newbery (*Trans. Electrochem. Soc.*, 1935, 67, 223-231; discussion, 231-233).—See *Met. Abs.*, this vol., p. 163.—S. G.

**\*Voltaic Couples and Corrosion.** Oliver P. Watts (*Trans. Electrochem. Soc.*, 1935, 67, 235-251; discussion, 251-257).—See *Met. Abs.*, this vol., p. 163.—S. G.

**On the Possibility of a Standardization of Corrosion Results.** F. Tödt (*Aluminium*, 1935, 17, 392-393).—In the present state of knowledge on corrosion problems the only safe criterion of the relative corrosibility of metals is considered to be the loss in weight expressed as grm./m.<sup>2</sup>/day.—A. R. P.

**The Chemist as Professional Corrosion Expert.** Fritz Tödt (*Wasser*, 1934, 8, (II), 124-133; *C. Abs.*, 1935, 29, 4316).—Discusses the chemical properties of metals (as solubility of their oxides, &c.) which are important in the matter of corrosion.—S. G.

## V.—PROTECTION

(Other than Electrodeposition.)

(Continued from pp. 385-386.)

**Ferrous Metals and High Temperatures. Protected Cast Irons and Steels.** Am. Matagrin (*Industrie chimique*, 1935, 22, 333-337).—This final part of a serial article discusses the value of aluminium, nickel, and chromium as protective coatings for iron and steel destined for use at high temperatures. The various methods of applying these coatings are described.—C. E. H.

**†Tinplate: Some Fundamental Considerations.** W. E. Hoare (*Proc. Swansea Tech. Coll. Met. Soc.*, 1934; and *Tech. Publ. Internat. Tin Res. Develop. Council*, 1935, [A], No. 14, 1-16).—A discussion of recent work on the reaction between iron and tin, the iron-tin compound in tinplate, and its influence on the mechanical properties of tinplate, the influence of annealing temperature on the alloy layer, the mechanism of the coating of steel with tin, and the porosity of tinplate.—E. S. H.

**\*A Study of the Yellow Stain on Tinplates.** C. E. Beynon and C. J. Leadbeater (*Tech. Publ. Internat. Tin. Res. Develop. Council*, 1935, [D], No. 1, 1-11).—The yellow stain which sometimes forms on tinplate during storage is due to surface oxidation of the tin in accordance with the differential aeration principle; it can be produced artificially by anodic oxidation. Difficulties in lithographing yellow-stained sheets cannot be attributed to the stain, as such sheets print satisfactorily when stoved at 140° C. or washed in methylated spirits—treatments which remove the film of grease which is believed to be responsible for the printing difficulty. Yellow stain can be removed by cathodic treatment of the sheet in dilute sodium carbonate solution. Since this treatment also removes the grease, the sheets can then be printed without difficulty. (See also following abstract.)—E. S. H.

**Causes of Yellow Stains on Tinplate, and Methods of Prevention.** E. T. Richards and R. J. Snelling (*Illust. Z. Blechindustrie*, 1935, 64, 887-888).—See also preceding abstract. An analysis is given of the conditions in which yellow stains are produced; the principal contributory factors are moisture, ventilation, temperature, and the quality of the tin employed. Storage in piles should be avoided. The staining is attributed to electrolytic oxidation, and it has been shown that it can be removed by appropriate electrical treatment. Preventive methods are summarized.—P. M. C. R.

**\*The Determination of the Porosity of Tin Coatings on Steel.** D. J. Macnaughtan, S. G. Clarke, and J. C. Prytherch (*Tech. Publ. Internat. Tin Res. Develop. Council*, 1935, [A], No. 7, 1-9).—Practical details are given of the determination of the porosity of tin coatings on steel by the hot-water method and the modified potassium ferrocyanide paper test. Improved methods for degreasing the specimens before applying the tests are described. The specimen is immersed in boiling 1% water-glass solution for 1 minute and then washed in water. Alternatively, the specimen is made a cathode in 0.5% sodium carbonate solution, using a sheet of nickel on each side as anodes. An e.m.f. of 4 v. is applied for about 1 minute, until gas bubbles form freely on the surface. Exceptionally resistant grease films may be removed by washing with carbon tetrachloride, applying a thin coating of a cellulose-base varnish, and then degreasing by the cathodic method, whereupon the varnish film is detached and removes all the grease with it.—E. S. H.

**\*A Rapid Test of Thickness of Tin Coatings on Steel.** S. G. Clarke (*Analyst*, 1934, 59, 525-528; and *Tech. Publ. Internat. Tin Res. Develop. Council*, 1934, [A], No. 12, 4 pp.).—The loss of weight of the specimen on stripping the tin coating in a solution of antimony chloride in hydrochloric acid is determined. The attack on the basis steel is negligible. With hot-dipped tin coatings the underlying layer of tin-iron alloy is almost completely dissolved, and a correction for the iron constituent amounting to one-third of the alloyed tin is necessary. The determination may be carried out on a selected area by protecting the rest of the coating with varnish.—E. S. H.

**The Surface Testing of Tinplate.** A. Karsten (*Illust. Zeit. Blechindustrie*, 1935, 64, 820-822).—An apparatus is described and illustrated for the optical testing of tinplate. The apparatus may be used with or without a camera attachment, and a modification, adopted for the examination of the internal surfaces of cylinders, is stated to have proved valuable in investigating cases of corrosion of tinned containers by foodstuffs.—P. M. C. R.

**Tinplate Manufacture.** J. Zelander (*Mech. World*, 1935, 97, 637).—A flowsheet, which gives in condensed form the whole process of tinplate manufacture.—F. J.

**Possibility of Substituting Zinc for Tin in Coating Copper Wire.** V. P. Kibanov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (4), 74-81; *C. Abs.*, 1935, 29, 4316).—[In Russian.] The process of covering copper wire with zinc is similar to tinning, except that the temperature in the first case is 450° C. while in the second it is 290°-310° C. The zinc coating is plastic and does not crack or peel on twisting the wire. It increases resistance 2.6-4.2%, depending on the size of wire, the increase being smaller for heavier wires. The zinc-covered wire, on vulcanizing, becomes coated with a thin layer of zinc sulphide, which, however, has no harmful effect on the wire or the rubber. The difficulties involved in the zinc coating process are oxidation of the liquid metal, in spite of the carbon protection, rapid cooling which does not permit the excess zinc to run off the wire, and the high temperature of the operation. The latter might be overcome by the addition of other metals.—S. G.

**Zinc Galvanizing. Effect of Impurities and Factors Governing Results with Various Types of Zinc.** Edmund R. Thews (*Canad. Chem. and Met.*, 1935, 19, 137-138, 142).—The most detrimental impurity in zinc for galvaniz-

ing is arsenic, since it reduces the fluidity of the bath, results in the heavy formation of ashes, and decreases the ductility of the coating; addition of aluminium eliminates to a greater extent the evil effects of arsenic, but in no case should the arsenic content exceed 0.01%, an amount which requires 0.03% of aluminium. Antimony in amounts not exceeding 0.01% produces a brown coloration on the coating, but about 1 oz. of aluminium per ton of zinc prevents this action. Copper should not exceed 0.05%, and iron should be kept to a minimum; lead is really beneficial and up to 1.5% may be tolerated, any excess above this figure settling to the bottom of the pot.

—A. R. P.

**How Far Below the Top of the Galvanizing Pot Should the Surface of the Galvanizing Bath Be?** Wallace G. Imhoff (*Amer. Metal Market*, 1935, 42, (134), 3, 6).—The best operating conditions as regards depth of metal and flux in the galvanizing pot are discussed with reference to the type of articles being galvanized. Heavy articles require a thick flux layer and small or light articles only a very thin layer or none at all; in both cases it is advisable to have the pot as full as possible to avoid crust formation and to assist in the maintenance of an equable temperature.—A. R. P.

**Flux on Hot-Dip Galvanizing Baths.** Wallace G. Imhoff (*Amer. Metal Market*, 1935, 42, (125), 6, 8).—A general discussion of the types of fluxes used on galvanizing baths, their functions and properties.—A. R. P.

**Bend Tests on the Hot-Dipped Zinc Coatings Measured with Micrometer Calipers.** Wallace G. Imhoff (*Amer. Metal Market*, 1930, 37, (247), 5, 9, 10).—The operating conditions of 4 American galvanizing plants are critically analyzed. Tabulation of the results shows that the amount of zinc deposited is a function of the time of immersion in and speed of withdrawal from the bath. When adherence is the important factor a light coating is best; ductile coatings are obtained by keeping the zinc as pure as possible, the bath well dressed and a steady production through the bath to maintain an even temperature therein.—A. R. P.

**Metallizing an Important Factor in Prevention of Corrosion in Refineries.** H. R. Leland (*Oil and Gas J.*, 1935, 33, (45), 99, 168).—S. G.

**The Leafing of Aluminium Paint.** P. H. Faucett (*Paint Varnish Production Mgr.*, 1935, 12, (May), 12; *C. Abs.*, 1935, 29, 4605).—Microscopic observation of the drying of the aluminium paint indicates that a circular cyclonic movement is set up in the film because of the escape of the volatile solvents. This causes the flat aluminium particles to align themselves parallel to the direction of movement of the liquid and in a position to offer the least resistance to the flow of the liquid past them. A viscous aluminium paint containing the optimum content of aluminium-bronze was taken and thinned with equivalent amounts of petroleum ether, petroleum benzine, V M and P naphtha, No. 1 distillate, and No. 2 distillate. The sample thinned with equal parts of these 4 liquids showed excellent leafing properties with resultant increase in hiding power of the product.—S. G.

## VI.—ELECTRODEPOSITION

(Continued from pp. 387-388.)

**\*Mechanism of Chromium Deposition from the Chromic Acid Bath.** Charles Kasper (*J. Res. Nat. Bur. Stand.*, 1935, 14, 693-709; and *Research Paper No. 797*).—In commercial electroplating chromium is deposited from baths composed principally of chromic acid in which the chromium is hexivalent. After use, such baths always contain some trivalent chromium, and many of the theories which have been proposed for this process postulate that the chromium is deposited from the trivalent (or possibly bivalent) state, for

which the chromic acid serves as a reservoir. K. presents evidence that the electrolytic deposition of chromium occurs directly from the sexivalent state. The principal evidence is the fact that relatively large additions of compounds of metals such as zinc, nickel, cadmium, iron, and copper have no such effect on the behaviour of the chromic acid bath as would be expected if deposition occurred from the trivalent state. It is shown by cryoscopic and spectrophotometric data that the dichromates of these metals are strongly ionized. The low efficiency and high polarization in chromium deposition are accounted for, and certain paradoxes are explained. The possible directions for improvement are pointed out, but none appears practicable with existing knowledge. A *bibliography* of 40 references is appended.—S. G.

**\*Influence of Bath Temperature on Chromium Hardness.** R. J. Piersol (*Metal Cleaning and Finishing*, 1935, 7, 169–172; *C. Abs.*, 1935, 29, 4677).—With a bath containing 2.50 gm. chromium trioxide and 2.5 gm. sulphuric acid per litre and operated at a current density of 1 amp./in.<sup>2</sup>, it was impossible to obtain a hard chromium plate at temperatures above 38° C. At lower temperatures the hardness increased gradually to the lowest limit investigated (21° C.). At a current density of 2 amp./in.<sup>2</sup> the hardest plate was obtained at 43° C. and the effect of the temperature was exceedingly critical between 49° and 52° C., with a decrease in hardness of from 100 to 5 units. A bath containing 250 gm. chromium trioxide, 2 gm. sulphuric acid, and 3 gm. Cr<sup>III</sup> per litre, operated at 68° C., produced hard deposits which were also very satisfactory as regards texture and smoothness. At this temperature the range of current density which produced very hard plates was extremely wide and was of importance for the plating of articles of irregular shape. The present results suggest future plating based on high current density and bath temperature and extremely rapid rates of deposition of chromium for wear-resistance.

—S. G.

**Industrial Applications of Chromium Plating.** Jacques du Chatenet (*Métaux et Machines*, 1935, 19, 270–273).—Comparative tests on special steels, nitrided steel, and steel protected by chromium plating showed that the chromium-plated specimen possessed a very high resistance to wear and corrosion. It is suggested that more extended use should be made of thick chromium deposits, as ordinary steels so protected can advantageously be substituted for the more costly special steels. Optimum adhesion and texture of the protective layer are obtained by its deposition in the cold.

—P. M. C. R.

**\*Direct Electrolytic Production of Copper Sheets.** A. A. Bulakh (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (7), 101–115; *C. Abs.*, 1935, 29, 4268).—[In Russian.] A study was made of the optimum conditions for the electrolytic production of copper plate on a rotating drum. It was found that successful deposits can be obtained at a cathode current density of 500–5000 amp./m.<sup>2</sup> from a copper sulphate solution containing 150–200 gm. sulphuric acid per litre. Addition of colloids is not desirable. Interruptions of the current for periods of 0.5 minute had no harmful effect on the quality of the product. Plates up to 0.25 mm. thick can be produced without the use of special compression rollers; for a thickness of 0.25–0.50 mm. compression is desirable, and for thicker plates compression is necessary. (Cf. U.S. patent No. 1,952,762.)—S. G.

**The Electrodeposition of Copper, Nickel, and Zinc Alloys from Cyanide Solutions. II.—Discussion of the Results of Part I.** Charles L. Faust and Geo. H. Montillon (*Trans. Electrochem. Soc.*, 1935, 67, 281–297; discussion, 297–298).—See *Met. Abs.*, this vol., p. 300.—S. G.

**\*Electrodeposits of Nickel-Iron Alloys.** E. Raub and E. Walter (*Mitt. Forschungsinst. Edelmetalle*, 1935, 9, 17–21).—Cf. *Met. Abs.*, this vol., p. 237. Bright deposits of iron-nickel alloys containing 40–60% iron can be obtained

from solutions containing nickel sulphate or chloride, ferrous sulphate, magnesium sulphate, and citric acid at a  $p_H$  of 3-3.6. The higher the temperature the lower is the iron content of the deposits, and at 100° C. the iron decreases to 12-14%. High temperatures reduce pitting, as also does addition of "Nekal," a mixture of alkali alkylnaphthalenesulphonates used as a wetting agent in the textile industry. Deposits with a high iron content tend to develop hair-cracks and have little protective power against rusting when applied to an iron surface.—A. R. P.

**Faulty Silver Deposits. Influence of Weighting Cements and Sulphur on Silver Plating.** E. Raub and K. Bihlmaier (*Mitt. Forschungsinst. Edelmetalle*, 1935, 9, 29-35).—Hollow articles of thin metal are often weighted with organic cements containing resins, asphalt, and oils with an inorganic filler, usually calcium carbonate; this type of cement is attacked by the ordinary silver-plating bath and organic compounds pass into the solution and produce rough, dark, and usually porous deposits. Since there is no method of removing the objectionable constituents from the bath, it is recommended that such articles be plated before the cement filling is inserted. Particles of suspended silver sulphide in the plating bath produce nodular deposits; in this case the bath may be heated and allowed to rest for a day or two to allow the sulphide particles to coagulate and settle.—A. R. P.

**\*Cadmium-Zinc Alloy Plating from Acid Sulphate Solutions.** Colin G. Fink and C. B. F. Young (*Trans. Electrochem. Soc.*, 1935, 67, 311-336; discussion, 336-338).—See *Met. Abs.*, this vol., p. 299.—S. G.

**\*Studies in Zinc Electrodeposition: Deposition from Ammonium Sulphate Zinc Baths.** Raymond R. Rogers and Edgar Bloom, Jr. (*Trans. Electrochem. Soc.*, 1935, 67, 299-309).—See *Met. Abs.*, this vol., p. 167.—S. G.

**\*A Further Study of Cyanide Zinc Plating Baths Using Al-Hg-Zn Anodes.** A. Kenneth Graham (*Trans. Electrochem. Soc.*, 1935, 67, 269-279; discussion, 279).—See *Met. Abs.*, this vol., p. 167.—S. G.

**New Plating Anode of Zinc, Aluminium, and Mercury.** — (*Amer. Metal Market*, 1935, 42, (91), 6).—A brief description is given of the advantages of using zinc containing small amounts of aluminium and mercury as an anode in zinc cyanide plating baths (cf. preceding abstract).—A. R. P.

**Electrodeposits on Aluminium.** G. Elssner (*Aluminium*, 1935, 17, 376-378).—A brief account is given of methods of preparing the metal for plating, of the application of anchoring undercoats, and of the final plating process.

—A. R. P.

**Technological Developments in the Plating Industry.** C. B. F. Young (*Metal Cleaning and Finishing*, 1935, 7, 165-168; *C. Abs.*, 1935, 29, 4269).—A discussion of the electrodeposition of zinc on steel wires, the electrolytic treatment of zinc or zinc-coated articles which greatly increases the resistance to corrosion and imparts a deep black colour to the zinc, and the allowance to be made in the thickness of copper deposits on zinc to compensate for the diffusion of zinc and copper during the plating. The electrodeposition of tin and indium are also briefly described.—S. G.

**\*The Use of a Modified Haring Cell in Detecting Electrode Reactions.** Jesse E. Stareck and Robert Taft (*Trans. Electrochem. Soc.*, 1935, 67, 357-374).—With the aid of a modified Haring cell, the deposition of silver from silver nitrate and from cyanide solutions was carefully studied. Starting with extremely low current densities and voltages, these were gradually increased and every slight break in the volt-ampère curves repeatedly checked. Finally, these breaks in the curves are interpreted and assigned to certain physical and/or chemical reactions. The cathode products detected were crystalline silver, silver hydride, and hydrogen; the anode products were silver peroxide and oxygen. The characteristics and properties of silver hydride formed at the cathode are discussed at length.—S. G.

## VII.—ELECTROMETALLURGY AND ELECTROCHEMISTRY

(Other than Electrodeposition.)

(Continued from p. 388.)

\***An Electrochemical Investigation of Solid Cadmium-Antimony Alloys.** Arne Ölander (*Z. physikal. Chem.*, 1935, [A], 173, 284-294).—The electrode potential of cadmium-antimony alloys was determined at 240°-290° C.; the results indicate that the stable intermediate phase CdSb has a range of homogeneity of 50-50.5 atomic-% cadmium. The thermodynamic relations and the degree of lattice disorder of the phase have been calculated. A table is given showing the statistical term of the temperature coeff. of the potentials with different degrees of lattice disorder.—B. Bl.

\***The Hydrogen Overvoltage and the Anodic Behaviour of Tungsten in Aqueous Solutions of Potassium Hydroxide.** M. de Kay Thompson and C. W. Rice, Jr. (*Trans. Electrochem. Soc.*, 1935, 67, 71-78; discussion, 79).—See *Met. Abs.*, this vol., p. 169.—S. G.

\***The Mechanism of Conductance.** Hiram S. Lukens (*Trans. Electrochem. Soc.*, 1935, 67, 29-36).—Presidential Address. See *Met. Abs.*, this vol., p. 351.—S. G.

## VIII.—REFINING

(Continued from p. 388.)

\***Use of Pure Oxygen in Lead Refining.** D. M. Chizhikov and M. N. Sokolov (*Zvetye Metally (Non-Ferrous Metals)*, 1934, (7), 59-73; *C. Abs.*, 1935, 29, 4303).—[In Russian.] Oxygen was blown through molten impure lead, contained in an iron crucible at lower temperatures and in a clay crucible at higher temperatures, by means of a porcelain tube. At 350°-500° C. a considerable time is required to remove all the zinc, arsenic, antimony, and tin, which leads to the formation of litharge (40-47% by weight of charge). At 860°-900° C. the removal of the impurities is almost complete in 5-10 minutes, and litharge formation is only 4.0-5.5%. As compared with air, oxygen speeds up the reaction and, therefore, results in a saving of fuel.—S. G.

## IX.—ANALYSIS

(Continued from pp. 388-391.)

**Determination of the Fineness of Silver Alloys by Streak Tests.** Karl Bihlmaier (*Mitt. Forschungsinst. Edelmetalle*, 1935, 9, 22-24).—Details are given of the tests using standard needles and a special  $\text{HNO}_3\text{-H}_2\text{CrO}_4$  acid mixture. The method is applicable only to Cu-Ag alloys, and with practice is accurate to  $\pm 2\%$  for alloys containing 60-100% Ag.—A. R. P.

**A Very Sensitive Reagent for Aluminium. Separation of Aluminium and Beryllium.** T. Gaspar y Arnal (*Anal. Soc. españ. Fis. Quím.*, 1934, 32, 866-871).—The reagent consists of a solution of 20 gm. of  $\text{Ca}_2\text{Fe}(\text{CN})_6 \cdot 12\text{H}_2\text{O}$  in 670 c.c. of water to which is added 400 c.c. of 96%  $\text{C}_2\text{H}_5\text{OH}$ . As little as 0.02 mg. of Al in 1 c.c. produces an opalescence. For gravimetric work the precipitation is carried out in a boiling solution, and the precipitate is dried at 85°-90° C. for weighing; alternatively a known amount of reagent may be added and the excess determined by titration with  $\text{KMnO}_4$  after removing the precipitate. Be gives a precipitate under the same conditions as Al but the Be compound dissolves on addition of water whereas the Al compound remains insoluble.—A. R. P.

**The Microscopic Identification of the Elements of the Platinum Group and Gold.** Willet F. Whitmore and Herman Schneider (*Mikrochemie*, 1935, 17, 279-319).—[In English.] Au is extracted from the solution by shaking with  $\text{CH}_3\text{CO}_2\text{C}_2\text{H}_5$ ; treatment of the extract (after evaporation of the solvent) with caffeine affords golden hair-like crystals. The aqueous layer from the extraction is distilled with  $\text{HNO}_3$  in a micro-distillation apparatus to remove the Os, and the residual solution is divided into several portions. Pd is tested for by addition of HCl and dimethylglyoxime in the usual way. A second portion is treated with solid methylamine hydrochloride; a deep green ring round the circumference of the drop and the formation of green crystals as the solution evaporates indicates Ru, while an immediate red crystalline precipitate (octahedra) indicates Ir. To a third portion is added *m*-toluidine hydrochloride; sheaves of yellow crystals indicate Pt. There is no known specific reaction for Rh. Tables are given showing the behaviour of the Pt group of metals and Au towards a large number of organic reagents.—A. R. P.

**\*The Determination of Trivalent Chromium in Chromic Acid and in Chromium Plating Baths.** Hobart H. Willard and Philena Young (*Trans. Electrochem. Soc.*, 1935, 67, 347-356; discussion, 356).—See *Met. Abs.*, this vol., p. 171.

—S. G.

**Quinaldinic Acid as a Microchemical Reagent. II.—Estimation of Copper and Its Separation from Cadmium and Manganese.** Priyadarajan Ray and Jagannath Gupta (*Mikrochemie*, 1935, 17, 14-16).—The method previously described (*Met. Abs.*, 1934, 1, 194) is modified slightly for microchemical work.

—A. R. P.

**\*Microchemical Determination of Gold in the Presence of Palladium and Tin.** J. Donau (*Mikrochemie*, 1935, 18, 11-17).—Alloys containing Pd and Sn should be given a double inquartation with the Zn-Cd eutectic alloy, the beads being parted first in 1 : 1 then in concentrated  $\text{HNO}_3$ . Decantation and washing of the ( $\text{SnO}_2 + \text{Au}$ ) residue are effected in a centrifuge and the  $\text{SnO}_2$  is removed by heating with  $\text{NH}_4\text{Cl}$  prior to weighing the Au.—A. R. P.

**\*The Determination of Metallic Magnesium, Magnesia, and Magnesium Fluoride in the Products of Electrolytic Recovery of Metallic Magnesium.** V. Ya. Tartakovskii (*Trans. Inst. Econ. Mineral. (U.S.S.R.)*, 1934, (64), 16 pp.; *C. Abs.*, 1935, 29, 4692).—[In Russian.] Two methods were developed for the analysis of crude electrolytic Mg containing  $\text{MgO}$  and  $\text{MgF}_2$ . These methods are based on the fact that, in the presence of  $\text{K}_2\text{Cr}_2\text{O}_7$  or  $\text{HF}$ , Mg becomes passive and insoluble in acids. In the first method the sample is treated with a mixture of  $\text{AcOH}$  and  $\text{K}_2\text{Cr}_2\text{O}_7$ ;  $\text{MgO}$  dissolves, and Mg and  $\text{MgF}_2$  are filtered off. The residue is then treated with dilute  $\text{AcOH}$ , which dissolves the Mg. In the second method,  $\text{MgO}$  is first converted into  $\text{MgF}_2$  by means of  $\text{HF}$ ; the Mg is then dissolved from the mixture of Mg and  $\text{MgF}_2$  with  $\text{AcOH}$ .  $\text{MgO}$  and  $\text{MgF}_2$  are determined by the first method.—S. G.

**Volumetric Determination of Tin Using Various Reducing Agents and Various Concentrations of Acid.** F. W. Louw and W. E. Schilz (*J. S. African Chem. Inst.*, 1934, 17, 3-7; *Chim. et Ind.*, 1934, 32, 791).—Fuse with  $\text{Na}_2\text{O}_2$ , dissolve in 200 c.c. hot water, and neutralize with HCl. Add 400 c.c. HCl, and then 1-2 gm. aluminium bromide. Pass a current of  $\text{CO}_2$  through the flask, and when the evolution of  $\text{H}_2$  ceases heat until ebullition just commences. On cooling, syphon over some  $\text{Na}_2\text{CO}_3$  solution, and titrate with  $\text{I}_2$  as usual. Other reducing agents used are: powdered Al (gives variable results), powdered Zn (good results), Ni foil, Pb foil (gives good but consistently low results). Mg cannot be used but Sb is satisfactory. Acid concentrations of 35-50% are the most satisfactory.—W. A. C. N.

**\*A Method for the Determination of Small Amounts of Zinc in Commercial Nickel.** B. S. Evans (*Analyst*, 1935, 60, 464-467).—The metal (10 grm.) is dissolved in  $\text{HNO}_3$  and the Fe removed by double precipitation with  $\text{NH}_4\text{OH}$ .



The filtrate is treated with KCN until it assumes a clear orange colour, then with  $\text{CH}_3\text{-CO}_2\text{H}$  until acid to litmus, saturated with  $\text{H}_2\text{S}$  at above  $60^\circ\text{C}$ . under pressure, cooled and filtered. The precipitate of  $\text{ZnS}$ ,  $\text{CuS}$ , and  $\text{PbS}$  is dissolved in  $\text{HNO}_3$  and  $\text{H}_2\text{SO}_4$ , the Cu removed with  $\text{H}_2\text{S}$ , and the Pb with  $\text{Na}_2\text{S}$  after addition of citric acid,  $\text{NH}_4\text{OH}$ , and KCN, and finally pure  $\text{ZnS}$  is precipitated by acidifying with formic acid and passing  $\text{H}_2\text{S}$  as before. The  $\text{ZnS}$  is converted into  $\text{ZnSO}_4$  for weighing.—A. R. P.

**Quinaldinic Acid as a Microchemical Reagent. I.—Estimation of Zinc and Its Separation from Manganese.** Priyadarajan Rây and Mukul Kumar Bose (*Mikrochemie*, 1935, 17, 11–13).—The method previously described (*Met. Abs.*, 1934, 1, 194) is adapted to micro-analysis.—A. R. P.

**Quinaldinic Acid as a Microchemical Reagent. III.—Estimation of Zinc in the Presence of Iron, Aluminium, Uranium, Beryllium, and Titanium.** Priyadarajan Rây and Mukul Kumar Bose (*Mikrochemie*, 1935, 18, 89–91).—The precipitation is made in slightly ammoniacal tartrate solution and the excess of  $\text{NH}_3$  is removed by blowing a current of air over the surface of the solution at  $60^\circ\text{C}$ . before filtering.—A. R. P.

**On the Microchemical Determination of Zinc by Means of Anthranilic Acid.** Ch. Cimerman and P. Wenger (*Mikrochemie*, 1935, 18, 53–57).—[In French.] The method of Funk and Ditt (*Met. Abs. (J. Inst. Metals)*, 1933, 53, 453) is modified for microchemical work.—A. R. P.

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## X.—LABORATORY APPARATUS, INSTRUMENTS, &c.

(See also “Testing” and “Temperature Measurement and Control.”)

(Continued from pp. 391–392.)

**\*An Induction Furnace in which Motion of the Bath is Eliminated, e.g. For the Melting of Light Metals.** E. Fr. Russ (*Z. Physik*, 1935, 93, 220–226).—Devices for reducing or eliminating motion of the liquid metal bath in induction furnaces are suggested. Motion of the bath liquid is attributed to divergence of the current stream lines from parallelism and the devices proposed eliminate this divergence. The elimination of motion of the bath in the case of aluminium results in castings being produced which are gas free, and very little metal is lost by burning.—J. S. G. T.

**\*Furnace Temperature and Heating-Rate Control by Dilating Wire.** C. T. Eddy, F. W. Vigelius, and E. C. Davis (*Instruments*, 1935, 8, 174–176).—Typical current and temperature curves are given to show that the temperature of a small laboratory resistance furnace can be varied at any desired constant rate by choosing a suitable starting current and increasing this uniformly. An arrangement for reproducing current (and thus temperature) runs consists of a control wire, mounted under tension in a glass tube, and connected in series with the furnace. The dilation of this wire controls a power-driven rheostat. Normally, the wire is kept at constant length, but by rotating a commutator the length of the wire, and thus the current passing, can be uniformly varied. Fluctuations in room temperature have no appreciable effect on performance.—J. C. C.

**An Intense Glow-Discharge Tube for the Spectroscopic Investigation of Very Small Masses of Material.** H. Schüler and H. Gollnow (*Z. Physik*, 1935, 93, 611–619).—A glow-discharge tube, suitable for the spectroscopic investigation of extremely minute samples, is described. The apparatus has already been applied to the detection of protactinium, cassiopeium (lutecium), terbium, samarium, scandium, thulium, yttrium, rhodium, holmium, and europium.—J. S. G. T.

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## XI.—PHYSICAL AND MECHANICAL TESTING, INSPECTION, AND RADIOLOGY

(Continued from p. 392.)

**Examination of Polished Surfaces.** J. M. Waldram (*Illuminating Engineer (Lond.)*, 1934, **27**, 163-164; *Sci. Abs.*, 1934, [B], **37**, 427).—The examination of a polished surface for the detection of minute flaws is generally carried out by observing the reflection of a luminous object in the surface. The flaws cause local distortion of the image, but in order that this may be noticeable the source must have a "structure," i.e. it must be of some extent but not uniformly bright. W. describes a specially designed source consisting of an extended bright surface crossed by a number of dark strips. The effect of flaws in causing deformation of the image of such a source is shown by means of photography.—S. G.

**Working Stresses.** C. Richard Soderberg (*Amer. Soc. Mech. Eng. Advance Copy*, 1932, Dec., 9 pp.).—The fundamental aspects of the problem of the strength and safety of machine parts is reviewed mathematically with especial reference to the behaviour of stainless and other special steels at high temperatures and under torsional, tensile, and shear stresses.—A. R. P.

**Open Discussion on the Metallurgical Inspection of Engineering Materials.**—(*Proc. Staff. Iron. Steel Inst.*, 1933-1934, **49**, 66-72).—See *Met. Abs.*, 1934, **1**, 254.—L. A. O.

**Commercial Testing of Engineering Materials.** I.—[Tensile, Compression, and Transverse Tests]. II.—Ductility Tests on Sheet Metals. III.—Impact Tests. IV.—Torsion Tests. V.—Hardness Tests. J. Trickett (*Mech. World*, 1935, **97**, 603-606, 631; 1935, **98**, 3-4, 29-30, 55-60).—Describes and illustrates: (I.—) The tensile test-piece, the tensile testing-machine, making the tensile test, compression and transverse tests. (II.—) The Erichsen cupping test and the Olsen ductility test. (III.—) The Izod and Charpy impact tests (of notched bars). (IV.—) Typical methods and machines for making torsion tests on metals. (V.—) The Brinell ball hardness test.—F. J.

**\*Determination of the Alternating Bending Strength of Wires. Construction of a Suitable Testing Machine.** Werner Friedmann (*Mitt. Wohler-Inst.*, 1934, (22), 91 pp.).—See also *Met. Abs.*, this vol., p. 174. To avoid fracture of the specimens in the holders in the Föppl-Heydekampf alternating bending machine the use of lead supports in the grips is recommended; even with this modification hard-drawn aluminium specimens frequently fracture in the holders. A modification of the machine for testing wires up to 3 mm. in diameter is described and the results obtained with it in steel, aluminium, Aldrey, copper, and various bronze wires in the annealed and hard-drawn states are tabulated. A selection of the results is given below for wires of 2.1-2.8 mm. diameter:

	Tensile Strength (A), kg./cm. <sup>2</sup> .	Modulus of Elasticity, kg./cm. <sup>2</sup> .	Alternating Bending Strength (B) (2 × 10 <sup>6</sup> reversals), kg./cm. <sup>2</sup> .	Ratio: B/A.
Soft copper . . .	2300-2400	1.3 × 10 <sup>6</sup>	1235-1300	0.54
Hard copper . . .	4500	1.3 × 10 <sup>6</sup>	1295	0.29
Hard aluminium (99.5%) . . .	1500-2000	0.7 × 10 <sup>6</sup>	590-920	0.34-0.53
Aldrey . . .	3000-3300	0.65 × 10 <sup>6</sup>	1060-1250	0.34-0.40
Cadmium-bronze . . .	6100-6300	1.34 × 10 <sup>6</sup>	1656-2020	0.27-0.33
Tin-bronze . . .	5800-6600	1.3 × 10 <sup>6</sup>	1370-2120	0.23-0.32

—A. R. P.

†**Materials Testing Machines.** C. H. Gibbons (*Baldwin Locomotives*, 1935, 13, (3), 31-38; 14, (1), 31-39; and *Instruments*, 1935, 8, 48-49, 76-78, 99-102, 130-132, 156-159).—See *Met. Abs.*, this vol., pp. 28, 111. This very comprehensive review of tension-compression-transverse testing machines is concluded. A *bibliography* of 79 references is contained in foot-notes. A tabulated comparison of the design features, accuracy, and sensitivity of the principal modern types of testing machines concludes the series as printed in *Baldwin Locomotives*, but is omitted from the reprint in *Instruments*.—J. C. C.

\***Endurance Bending Strength of Constructional Parts at Sockets, Boss-Heads, and Similar Joined Parts.** A. Thum and F. Wunderlich (*Mit. Materialprüfungsanst. Tech. Hochschule Darmstadt*, 1934, (5), 82 pp.).—Endurance tests on flat and round rods of various metals with one end inserted in various shaped sockets and bosses have been made under vibrational stresses to ascertain the effect of such joints on the strength of complex structures. The endurance strength decreases with the pressure in the socket and tends to reach a limiting value depending on the nature of the 2 metals in contact, which is lower the more sensitive is the rod to notch effects, and higher the softer the metal of the socket. Oil in the junction raises the endurance strength, but graphite has no effect and water tends to reduce it. The effects of surface condition, case-hardening, fractional wear, &c., are considered chiefly with reference to ferrous metals. Numerous practical examples are discussed in detail.—A. R. P.

\***Modulus of Elasticity.** Léon Guillet, Jr. (*Rev. Mét.*, 1935, 32, 61-68).—A study of the modulus of elasticity in tension and bending. Methods of determination are discussed. Attention is directed to the important effect of temperature, a variation of 5° C. resulting in an error of the order of 5%. The practical advantages and theoretical disadvantages of the "bending" method of determining Young's modulus are detailed. The dynamic method of Rolland and Sorin is simple, rapid, and gives results in good agreement with direct observations obtained from careful extensometer tests, results of both types of tests being given. The influence of graphite on the modulus of cast iron is considered. Modulus values for the different elements plotted against melting temperature divided by the square of the atomic volume results in fair alignment of the points. G. discusses the influence of chemical composition and treatments on modulus value and gives a short *bibliography*.—H. S.

**Tensile Testing Machine, 2.75 kg. for Small Test-Pieces.** R. Guillery (*Rev. Mét.*, 1935, 32, 58-60).—A small hydraulic machine of the bench type.—H. S.

\***New Machine for Hardness Tests of Metals.** R. Guillery (*Rev. Mét.*, 1935, 32, 49-53).—The preliminary load on the ball is  $\frac{1}{5}$  of the main load subsequently applied. A steel ball is used for tests of soft metals and tungsten carbide for tests of hard steels.—H. S.

#### RADIOLOGY

\***Radiography of Fine Flaws in Metals.** A. G. Warren (*Brit. J. Radiology*, 1935, 8, 235-247).—Cracks and crack-like voids present the most difficult problem in the radiographic examination of metal structures. In order to determine the practical limits of examination artificial cracks in steel have been examined. Examples of typical cracks found in practice are given and a method of radiography applicable to cylindrical objects is mentioned.

—S. V. W.

**Inspection of Welds—With Particular Reference to Radiography.** Herbert R. Isenburger (*Welding Eng.*, 1935, 20, (6), 26-27).—The advantages of radiographic inspection are pointed out, some hints on the choice of suitable equipment are given, and the importance of exact technique and correct interpretation is emphasized.—H. W. G. H.

†The Application of Very Hard X-Rays to the Testing of Materials. A. Herr (*Elektrotech. Z.*, 1935, 56, 425-428).—Reasons are given for the use of very hard X-rays in the testing of metals and some of their technical applications are discussed with reference to examples. A description of the necessary apparatus and of the precautions to be taken in its use is also given.—B. Bl.

## XII.—TEMPERATURE MEASUREMENT AND CONTROL

(Continued from p. 393.)

Temperature Indicator Methods in Aluminium Switchgear. C. Geier, W. von Zwehl, and W. Helling (*Aluminium*, 1935, 17, 375-376).—The various coloured compounds, chiefly derivatives of mercuric and other iodides, usually employed for indicating dangerous increases in temperature by a colour change can be employed on aluminium switchgear if the metal is first protected by a lacquer coat.—A. R. P.

Realization of the International Temperature Scale by Means of Thermocouples of Platinum and Platinum-Rhodium. V. N. Bozhovskii and A. S. Arzhanov (*Vsesoyuznuii Nauch.-Issledovatel. Inst. Metrol. i Standartizatsii*, 1933, (132), 3-11 (in French), 29-33 (in Russian); *C. Abs.*, 1935, 29, 4641).—A comparison over the range 300°-1300° C. of the International Temperature Scale as determined by the melting points of gold, silver, and antimony, with a scale determined by the melting points of zinc, antimony, and copper.—S. G.

Realization of the International Temperature Scale in the Interval 0° to 600° C. by Means of the Platinum Resistance Thermometer. V. A. Yakovleva (*Vsesoyuznuii Nauch.-Issledovatel. Inst. Metrol. i Standartizatsii*, 1933, (132), 22-28 (in French), 41-44 (in Russian); *C. Abs.*, 1935, 29, 4642).—A determination of the melting point of antimony is included.—S. G.

Realization of the Fundamental Reference Point of Optical Pyrometry. V. N. Bozhovskii and V. I. Parvitzkii (*Vsesoyuznuii Nauch.-Issledovatel. Inst. Metrol. i Standartizatsii*, 1933, (132), 12-21 (in French), 34-40 (in Russian); *C. Abs.*, 1935, 29, 4641).—Apparatus and method are described.—S. G.

## XIII.—FOUNDRY PRACTICE AND APPLIANCES

(Continued from pp. 393-395.)

Removal of Zinc from High-Tin Bronze. N. N. Muratch (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (4), 69-74; *C. Abs.*, 1935, 29, 4316).—[In Russian.] By blowing air through molten bronze scrap the zinc in it was reduced to the permissible limit. With molten scrap containing zinc 1.09, tin 18.27, copper 79.19, iron 0.03, lead 0.92, and nickel 0.10%, the final product, after blowing with air for 6 hrs. 20 minutes and subsequent addition of cupric oxide to the slag, contained: zinc 0.08, tin 12.78, copper 85.70, iron 0.04, lead 0.92, and nickel 0.15%.—S. G.

Foundry Production of Nickel Silver.—II. T. E. Kihlgren (*Metals and Alloys*, 1935, 6, 175-180).—The choice of a suitable deoxidizer for nickel-brass castings is discussed. Silicon in the alloy tends to produce porous brittle castings, hence deoxidizers, such as lithium, boron, and titanium which reduce silica from the melting crucible, should not be used. The best results are obtained by the use of phosphorus, manganese, and magnesium as described in section I (*Met. Abs.*, this vol., p. 394). Numerous examples and illustrations showing the effects of various deoxidizers and correct methods of moulding and casting are given.—A. R. P.

**Foundry Practice for Nickel Silver (Including Some Other Nickel-Containing Non-Ferrous Alloys).** James A. Rabbitt (*Japan Nickel Information Bureau Working Instructions*, C-16, 1935, 106 pp.).—[In English and Japanese.] The quality of raw materials, melting equipment and practice, and moulding technique are discussed. Several alternative procedures for melting nickel-brass are described, but the recommended method is that specified by the Japanese Navy, in which a 50:50 copper-nickel alloy is first made and ingotted. Particulars of permissible impurities in nickel-brass castings, and typical charge sheets for nickel-bronzes, nickel-aluminium-bronzes, and light alloys are given. A list of typical nickel-brasses, with their trade names and compositions forms an appendix.—H. W. G. H.

**\*Removal of Sulphur from Tin and Its Alloys.** A. A. Bulakh (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (8), 115-116; *C. Abs.*, 1935, 29, 4721).—[In Russian.] Tin (285 kg.) containing sulphur 0.2 and iron 0.61% was heated to 530° C. and sodium hydroxide was added in the following manner: first, 5 kg. was added, after 25 minutes the slag was removed, 5 minutes later 3 kg. sodium hydroxide was added and after 25 minutes the slag was removed. The resulting metal contained a trace of sulphur and 0.11% iron.—S. G.

**\*Melting Zinc Dust under Fluxes.** N. N. Muratch and G. K. Markarov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (2), 107-108; *C. Abs.*, 1935, 29, 4304).—[In Russian.]  $\text{NH}_4\text{Cl}$  and  $\text{MCl}_2 + \text{NaHSO}_4$ , where  $M$  is alkali or alkaline earth, were tested as fluxes for zinc dust. The reactions are:  $\text{ZnO} + 2\text{NH}_4\text{Cl} = \text{ZnCl}_2 + 2\text{NH}_3 + \text{H}_2\text{O}$  and  $\text{ZnO} + \text{MCl}_2 + 2\text{NaHSO}_4 = \text{ZnCl}_2 + \text{Na}_2\text{SO}_4 + \text{MSO}_4 + \text{H}_2\text{O}$ . Both fluxes proved successful.—S. G.

**Effect of Rapid Cooling on the Distribution of Lead in Cast Zinc.** M. D. Zudin (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (5), 115-120; *C. Abs.*, 1935, 29, 4303).—[In Russian.] Rapid cooling of zinc ingots results in a finer crystal grain and more uniform distribution of lead.—S. G.

**Joint Plates and Joint Grids.** J. W. Molder (*Iron Steel Ind.*, 1935, 8, 421-422).—The use of joint plates and joint grids in the making of large castings for which special boxes are not available are discussed, and the making of both plates and grids and their operation are described in detail.—J. W. D.

**Die-Casting with Machines of Simple Design.** Charles O. Herb (*Machinery (N.Y.)*, 1935, 41, 669-672).—Hand-actuated semi-automatic die-casting machines can be used economically for making small parts up to 1 lb. in weight at a rate of 4 to 10 "shots" per minute. Descriptions are given of 2 dies for use with such machines, one casting 8 lead battery bushings and the other a zinc bowl-shaped part.—J. C. C.

**Die-Cast Wheel Hubs and Caps.** E. Stevan (*Mech. World*, 1935, 97, 5-6).—An example is given of economies effected by substituting die-cast hubs and caps for parts previously made from pressings (hubs) and brass sand-castings (caps). The details of the mould used are illustrated and described, but the composition of the die-casting alloy is not given.—F. J.

**†America Appreciates Zinc Die-Castings! And Germany? W. Mueller (*Maschinenbau*, 1935, 14, 67-72).**—The present position of the die-casting industry is discussed and the use of zinc die-castings in place of more expensive metals is reviewed in the light of modern metallurgical knowledge.—K. S.

#### XIV.—SECONDARY METALS: SCRAP, RESIDUES, &c.

(Continued from p. 395.)

**\*Treatment of Brass Scrap and Crude (80-90%) Copper at the Moscow Molotov Plant.** G. A. Shakhov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (5), 100-114; *C. Abs.*, 1935, 29, 4301).—[In Russian.] Cf. *Met. Abs.*, 1934, 1, 457. A study of the operation of tilting open-hearth furnaces and Bes-

semer converters for the refining of copper from brass scrap and crude copper, at the Moscow Molotov plant, led to the following conclusions: (1) the cost of refining in the converters is only  $\frac{1}{2}$  of that in the open-hearth; (2) one converter can produce as much as 3 open-hearth furnaces; (3) removal of antimony is complete in the converter and zero in the open-hearth; (4) consumption of coke in the converter is 10% by weight of that in the open-hearth; crude oil 30% by weight; (5) recovery of zinc in the converter is greater than in the open-hearth; (6) conditions of work at the converters are much better than at the open-hearth.—S. G.

**Extraction of Zinc from Waste Materials at the Belovo Zinc Plant.** V. F. Veber (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (9), 14–22; *C. Abs.*, 1935, 29, 4711).—[In Russian.] A description of methods used in the recovery of zinc scrap, dust, and other waste materials.—S. G.

**Industrial Non-Ferrous Metal Scrap. Its Classification and Preparation.** E. G. Wertheimer (*Amer. Metal Market*, 1935, 42, (98), 5; (99), 4).—Practical hints are given for sorting and classifying scrap metal from foundries and workshops with the object of accumulating uniform parcels which are more readily saleable and fetch better prices than do mixed parcels.—A. R. P.

**Avoiding Evils in Scrap Transactions.** Richard V. Bonomo (*Amer. Metal Market*, 1935, 42, (99), 9).—A brief discussion of the best methods for disposing of scrap metals in the U.S.A.—A. R. P.

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## XV.—FURNACES AND FUELS

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(Continued from p. 395.)

**Heat-Treating Aluminium Alloys.** L. H. Knapp (*Elect. World*, 1935, 105, 1712, 1715).—A brief, illustrated account of a sodium nitrate bath, employing 19,000 lb. of the salt, heated by twenty-four 7-kw. cast-in immersion heaters, operating at  $940^{\circ} \pm 3^{\circ}$  F., having a container made of  $1\frac{1}{8}$  in. welded boiler plate, and used for heat-treating aluminium alloy aircraft parts.—J. C. C.

**Heating of Furnaces.** — Guillon (*Usine*, 1935, 44, (18), 31; *C. Abs.*, 1935, 29, 4638).—The generation of heat in the furnace and its transmission to the material to be heated is discussed as dependent on construction, type of fuel and material.—S. G.

†**Air-Heating Blowers.** E. Schirm (*Feuerungstechnik*, 1935, 23, 61–64).—Blowers, incorporating a form of heat-exchanging device for heating gases, more especially air, are described.—J. S. G. T.

**Technical Considerations Relating to Heat Conservation Concerned in the Construction of an Electric Furnace.** Otto Krebs (*Feuerungstechnik*, 1935, 23, 54–55).—Thermal data relating to heat distribution, losses, &c., in an electric furnace are discussed.—J. S. G. T.

**Electric Heat for the Printer.** W. E. Swale (*Elect. Rev.*, 1935, 117, 80).—Electric melting, with thermostatic control, is ideal for type metal in a printing works. Tests on an electrically heated monotype caster showed it to consume 1.693 kWh/hr. A 3 kw. remelting pot consumed 25.2 Wh. per lb. of metal.—J. C. C.

**The Reactance of Large Rectangular Three-Phase Electric Furnaces.** F. V. Andreae (*Trans. Electrochem. Soc.*, 1935, 67, 151–167; discussion, 167–168).—See *Met. Abs.*, this vol., p. 359.—S. G.

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## XVI.—REFRACTORIES AND FURNACE MATERIALS

(Continued from p. 395.)

**\*Effect of Variations in the Alumina-Silica Ratio in Fused Refractory Materials upon Resistance to Metal Oxides [of Iron and Brass].** Charles McMullen, A. A. Turner, and G. J. Easter (*J. Amer. Ceram. Soc.*, 1935, 18, 30-33).—Solid pieces of electrically-fused material, varying in composition from alumina 63, silica 37% to 100% alumina, of the ordinary abrasive grade, were heated in contact with cast iron and brass which were allowed to oxidize and attack the refractory. The least attack was found in the case of iron with alumina 80 and silica 20%, and in the case of brass with straight alumina, although alumina 80:silica 20% was not greatly inferior. Bonds were developed for the materials thus selected and confirmatory tests were carried out on crucible fusions. Arc and powdered-coal fired furnaces lined with the bonded materials have proved successful.—S. G.

**Insulating Refractories.** W. M. Hepburn (*J. Amer. Ceram. Soc.*, 1935, 18, 13-17).—Insulating materials of various types are discussed, and their installation in industrial furnaces, &c. (including some metallurgical examples) is described. Data are given of savings involved in the use of insulating refractories. The characteristics of the materials are uniformity of product, shrinkage and spalling properties, physical strength, insulating value, and thermal capacity.—S. G.

**\*Physical Properties of Some Insulating Brick.** Walter C. Rueckel (*J. Amer. Ceram. Soc.*, 1935, 18, 18-22).—A study was made of 11 types of insulating brick. The physical properties determined included bulk density, porosity, transverse and crushing strength, pyrometric cone equivalent, volume shrinkage at 2300°, 2500°, and 2700° F., load deformation at elevated temperatures, and resistance to spalling. A study of the relation between permeability and pressure differential through the various insulating bricks is also reported. Suggestions are made for classifying the various types of insulating bricks into groups, and a discussion is given of formulæ for permeability.

—S. G.

**The Application of Insulating Materials to Industrial Furnaces.** J. D. Keller (*J. Amer. Ceram. Soc.*, 1934, 17, 77-87).—Deals with the classification of insulating materials, insulation applied outside refractory walls, the use of refractory insulators for the entire furnace wall, economics, and obstacles to the wider use of insulation.—S. G.

**Principles of Heat-Flow in Porous Insulators.** H. W. Russell (*J. Amer. Ceram. Soc.*, 1935, 18, 1-4; discussion, 4-5).—Gives an approximate theory of the thermal conductivity of porous materials. The effect of pore size and shape is discussed.—S. G.

**\*The Thermal Conductivity of Refractories.** Gordon B. Wilkes (*J. Amer. Ceram. Soc.*, 1934, 17, 173-177).—The thermal conductivity of recrystallized silicon carbide, clay-bonded silicon carbide, and silica and fireclay bricks was determined from 200° to 2600° F. (93°-1409° C.). The results are compared with those of previous investigators and a possible explanation is given of the peculiar variation of the coeff. of thermal conductivity with temperature for certain bricks.—S. G.

**\*A Method of Measuring Thermal Conductivity [of Refractory Materials] at Furnace Temperatures.** Clarence E. Weinland (*J. Amer. Ceram. Soc.*, 1934, 17, 194-202).—An apparatus of original design for the measurement of the thermal conductivities of insulating and refractory materials at temperatures up to 2600° F. (1409° C.) was constructed. The primary heat is supplied by Globar resistors, and the rate of heat-flow measured by a calibrated "heat-meter," means for the calibration of which were incorporated in the apparatus itself. The test-specimen consisted of 8 standard-size bricks form-

ing a section  $18 \times 18 \times 2\frac{1}{2}$  in. in size. Results are given of 6 complete tests: one of a high-temperature insulating brick, one of a fireclay brick, and 4 tests of refractory insulating bricks. In two cases comparative results from tests in a different kind of apparatus are shown. The accuracy of the results obtained in testing insulating and refractory insulating bricks are considered to be fully equal to the requirements of engineering use.—S. G.

**\*An Apparatus for Measuring the Thermal Conductivity of Refractories at High Temperatures.** J. L. Finck (*J. Amer. Ceram. Soc.*, 1935, 18, 6-12).—An apparatus is described for measuring the thermal conductivity of refractories up to about 2000° F. (1093° C.). The theory and method of test are discussed in detail and results are given for light refractory insulating brick and for a silica fire-brick and dense fire-brick.—S. G.

**\*A Study of the Effect of Natural Gas and of Hydrogen upon Various Refractories.** Bertram C. Ruprecht, R. H. H. Pierce, and Fred A. Harvey (*J. Amer. Ceram. Soc.*, 1934, 17, 185-193).—Natural gas was passed through an electric tube furnace where it was partially cracked and then through a second furnace containing the sample of a refractory. The refractories tested were fireclay bricks of various types and fired in various ways, and bricks made of magnesite, high alumina material, and silica. The action on the refractory was similar to that of carbon monoxide. Carbon was deposited in some of the refractory bodies within a critical temperature range. Some of the fireclay bodies were almost completely disintegrated by the growth of the carbon deposits, although others were scarcely affected. Hydrogen gas had no disintegrating action at any temperature in the range studied.

—S. G.

**The Destruction of Refractory Melting Furnace Brick-Work by Splintering.** E. T. Richards (*Feuerungstechnik*, 1935, 23, 49-52).—The splintering of refractory furnace materials due to mechanical and thermal effects and owing to textural changes in the material is discussed and illustrated.—J. S. G. T.

**\*A Rapid Method for the Analysis of Siliceous Refractories.** Sandford S. Cole and S. R. Scholes (*J. Amer. Ceram. Soc.*, 1934, 17, 134-136).—Evaporation with hydrofluoric and sulphuric acids removes silica. Ignition of the residue at 800°-900° C. yields a mixture of  $R_2O_3$  and sulphates, which is weighed. The residue is brought into solution and  $R_2O_3$ , calcium oxide, and magnesium oxide are determined by standard methods. Calcium and magnesium oxides are calculated back to sulphates and alkali is calculated from weight of ignited residue. Silica is found by difference. A summary is given of previous investigations on the behaviour of sulphates at elevated temperatures, as well as the results of similar experiments by C. and S.—S. G.

**A Compilation of Phase-Rule Diagrams of Interest to the Ceramist and Silicate Technologist.** F. P. Hall and Herbert Insley (*J. Amer. Ceram. Soc.*, 1933, 16, 463-567).—Gives a compilation of equilibrium diagrams, principally of silicates and refractory oxides. The text of the paper includes definitions of terms used in discussion of the Phase Rule, a description of methods used in the study of the systems at high temperatures, and an interpretation of Phase Rule diagrams for systems of one or more components.—S. G.

**The Physico-Chemical Study of the Manufacture of Refractory Materials.** M. Lépingle (*Métaux et Machines*, 1935, 19, 117-120, 257-262).—The Phase Rule is applied to the equilibrium diagrams of the principal types of refractories; the properties on which the refractory character of the material depends are thence deduced. Details of permissible variations in composition, working temperature range and reversible and irreversible thermal changes are given in each case. A table shows the influence of 26 catalysts on the transformation speed of quartz at 1300° C., and further tables show the effect of repeated heating.—P. M. C. R.



## XVII.—HEAT-TREATMENT

(Continued from p. 396.)

†**Bright-Annealing of Non-Ferrous Metals.** A. G. Robiette (*Metal Treatment*, 1935, 1, 63–69).—It is not sufficient to exclude oxygen to obtain a bright-anneal; water vapour, carbon dioxide, and gases evolved from the metal may all cause oxidation. The gas-metal equilibria, both at the annealing temperature and during cooling, must be taken into account, as well as such factors as the effect of a continuous gas stream in disturbing equilibria. Zinc alloys present troubles due to volatilization, and in addition will be oxidized if more than 0.2% of water vapour, carbon dioxide, or oxygen be present. By adding hydrocarbons to the gas mixture, however, oxide formation can be largely prevented. This action is apparently connected with the catalytic breakdown of the hydrocarbons. Nickel can be bright-annealed in the presence of water vapour and carbon dioxide provided sufficient reducing gases are present. Copper can be annealed in steam (wet annealing) or carbon dioxide, but traces of oxygen attack the metal above 80° C. Dry annealing is necessary for fine wires, and can be carried out without fear of "gassing" the copper if the atmosphere is only very slightly reducing. Lubricants on the surface may give trouble through carbon deposition or the liberation of free oxygen. The costs and applications of commercial atmospheres are reviewed, and the principal features of the main types of continuous and batch bright-annealing furnaces described.

—J. C. C.

**Practical Data Regarding the Heat-Treatment and Workability of Light Metals [Duralumin].** B. Adaridi (*Tek. Tids. Uppl. C., Bergsvetenskap.*, 1935, 65, 21–24, 31–32; *C. Abs.*, 1935, 29, 4307).—The effect of various heat-treatments on the tensile strength, elasticity, and elongation of rolled and drawn Duralumin is described, together with practical information on the working of Duralumin by hand-forming, cold-pressing after quenching in water, pressing in the annealed condition, and hot-pressing.—S. G.

## XVIII.—WORKING

(Continued from p. 396.)

**Note on the Treatment of Light Alloys in U.S.S.R. The Forging, Drop Forging, and Heat-Treatment of Elektron.** Michel Précoul (*Chaleur et Industrie*, 1935, 16, 241–242).—Elektron, used for propeller blades in the U.S.S.R., should be heated to 380° C. for 12 hrs. before forging and worked with frequent turnings between 330° and 380° C., reheating as required. Working increases the strength, especially in the longitudinal direction, but reduces elongation. Heat-treatment for 8 hrs. at 350° C. equalizes the strength in different directions at 24–25 kg./mm.<sup>2</sup>, and increases the elongation to 9–15%.—J. C. C.

**Stamping Monel Metal.** V. M. Aristov (*Zvetye Metally (The Non-Ferrous Metals)*, 1934, (4), 81–86).—[In Russian.] A review.—S. G.

**The Henley Lead-Extrusion Machine.** — (*Electrician*, 1935, 115, 148; also *Elect. Times*, 1935, 88, 103; and *Elect. Rev.*, 1935, 117, 118).—A continuous cable-sheath extrusion press has been perfected by W. T. Henley, Ltd., using the principle of a screw extruder. The machine is illustrated and briefly described. Its length from back to front is only about 12 in., and on account of the rigid construction, extremely close dimensional limits can be maintained in the extruded sheath. The usual oxide inclusions and seams are eliminated by the continuous nature of the extrusion.—J. C. C.

**Copper Wire Rope.** A. G. Arend (*Mech. World*, 1935, 97, 273-274).—Copper wire rope, although made in a smaller class of machine, with different number of strands, and in smaller quantity than steel rope, is used for haulage and transport in chemical factories and in the operation of all classes of tank plungers where corrosive fluids are used. Brass and iron wire ropes are too readily attacked in such conditions. Copper wire rope is not suited to withstand considerable abrasion, but when this property, together with the usual flexibility, is required, the wires in the outer layer are large, being laid on a layer of wires of small diameter. Details of the manufacture of copper wire rope are described. The progress recently made in radio and certain electrical pursuits has been responsible for an increased production.—F. J.

**Nichrome Wire.** M. W. Pridanzev (*Katschestvennaja Stalj*, 1934, 44-47; *Korrosion u. Metallschutz*, 1935, 11, 163).—Nichrome is made most readily by adding pieces of low-carbon nickel-chromium steel to molten 98% nickel and deoxidizing with ferrosilicon. The Russian alloy made in this way contains nickel 61.4, chromium 16.3, iron 19.5, carbon 0.12, silicon 0.5, manganese 2.4, phosphorus 0.012, and sulphur 0.007%. Wire made from this alloy is descaled after annealing by pickling at 40°-80° C. in a solution containing hydrochloric 5-20, nitric 5-20, and sulphuric acid 5-35%; it is then usually copper-plated.—A. R. P.

**\*Processes for the Production of Nichrome Wire.** B. F. Rukin (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (6), 81-100; *C. Abs.*, 1935, 29, 4307).—[In Russian.] A description of experiments for the production of Nichrome wire from Russian raw materials.—S. G.

**Free-Cutting Aluminium for Screw Machine Work.** — (*Amer. Metal Market*, 1935, 42, (131), 5, 6).—The free-cutting properties of aluminium are considerably improved by addition of lead and bismuth together with copper and magnesium. A special technique is required in the melting, casting, and working operations to obtain homogeneous distribution of the lead and bismuth which are insoluble in solid aluminium but soluble to a limited extent in the liquid metal at a high temperature. Age-hardening alloys of the Duralumin type may also be given free-cutting properties by the same process. The semi-hardened alloys work best in automatic lathes, but modifications in the shape of the tools are necessary; subsequently the screws may be hardened by heat-treatment. A general account of the development and properties of these alloys is given without any details of the composition or methods of working.—A. R. P.

**Working Aluminium and Its Alloys.** — (*Mech. World*, 1935, 97, 615, 619).—Technical details are given of the tools used for drilling, turning, milling, and sawing of aluminium and its alloys. On the whole, fine to medium feeds combined with high speeds, give the best results. It is preferable to use a cutting compound. A soluble cutting oil, or a mixture of lard oil and kerosene oil, the relative proportions varying according to the type of work and the speed and feed, will be found quite suitable. Pure lard oil is excellent for the heavier work.—F. J.

**The Need for Zinc Cutters.** — (*Mech. World*, 1935, 97, 517).—Letter to the Editor. The need for a method of cutting zinc sheets and cathode slabs by welding methods and the practical difficulties thereby involved are described.—F. J.

## XIX.—CLEANING AND FINISHING

(Continued from p. 397.)

**Electrolytic Method for Obtaining Bright Copper Surfaces.** P. A. Jacquet (*Nature*, 1935, 135, 1076).—The specimen is treated with emery paper, then cleaned cathodically in an alkaline solution, and finally treated anodically for

2 or 3 minutes in aqueous orthophosphoric acid (50 vol.-% of the commercial product, sp. gr. 1.71), using a high current density (about 60 amp. per sq. dm.). The solution tends to increase in temperature and should be cooled. A rotating anode is an advantage in securing uniformity.—E. S. H.

**Recent Advances in Metal Cleaning Technology.** R. W. Mitchell (*Metal Cleaning and Finishing*, 1935, 7, 133-135; *C. Abs.*, 1935, 29, 4307).—See also *Met. Abs.*, this vol., p. 360. A discussion of alkaline cleaners and new developments in cleaning equipment.—S. G.

**Chemical Stoneware in the [Cleaning] Finishing Industries.** Percy C. Kingsbury (*Metal Cleaning and Finishing*, 1935, 7, 239-242).—S. G.

**Tested Cleansers for Dairy Plants.** Forrest Grayson (*Food Industries (N. Y.)*, 1935, 7, 231-232, 281-282).—Sodium metasilicate is considered the best alkaline cleanser, as it has little action on tin, aluminium, and other metals, and a strong emulsifying and solvent action for milk fat. It is further improved by the addition of 5-10% of soap. Recommended concentrations for different pieces of plant are given.—C. E. H.

**Blackening Metals.** — Klement'ev (*Novosti Tekhniki, Seriya Gornorudnaya Promishlennosti*, 1935, (7), 9-10; *C. Abs.*, 1935, 29, 4307).—[In Russian.] The articles to be blackened are immersed in a saturated nitric acid solution of copper at 50°-60° C., removed, and heated until the liquid on the plate is evaporated and the greenish residue becomes black. The surface is then carefully brushed and the above operation repeated a number of times.—S. G.

**\*Atmospheres for Annealing High Brass in Relation to Buffing.** L. Kroll and E. A. Anderson (*Metals and Alloys*, 1935, 6, 192-193).—Experiments have been carried out on the loss in weight on buffing brass containing copper 65-75, lead 0.01, and iron 0.025%, and balance zinc. When the alloy is pickled in 7-15% sulphuric acid after annealing and then buffed the loss in weight increases with increase in annealing temperature, being about twice as great after annealing at 700° C. as after annealing at 440° C. At constant annealing temperature, increase in the time of annealing increases the buffing loss. Annealing in hydrogen produces smaller losses than annealing in air, but nitrogen atmosphere is not so good as air. Annealing in coal-gas is definitely unsatisfactory owing to the discoloration produced by the sulphur content of the gas and by carbon deposits derived from the cracking of the hydrocarbons.—A. R. P.

## XX.—JOINING

(Continued from pp. 397-401.)

**Contribution to the Investigation of Fusion Joints Between Aluminium and Other Metals.** — Holler and — Maier (*Autogene Metallbearbeitung*, 1935, 28, 177-187).—Joints with iron, copper, brass, nickel, Monel metal, stainless steel, lead, and zinc are described and illustrated by photomicrographs. Details of the technique used and physical properties obtained are given. In most cases, the heavy metal is "tinned" before welding. Corrosion tests in tap-water and 3% brine show that the joints have about the same resistance to corrosion as soft-soldered joints in aluminium.—H. W. G. H.

**Welded Light Metal Wagons of the French Northern Railway.** F. Thomas (*Aluminium*, 1935, 17, 378-380).—The welding technique is explained with reference to illustrations.—A. R. P.

**\*Welding of Zinc Sheets.** Hans A. Horn (*Autogene Metallbearbeitung*, 1935, 28, 169-171).—Samples up to 2.5 mm. thick were welded by oxy-acetylene, with a flame having about 10% excess oxygen. Flux-cored filler-rods, and also strips cut from the sheet with the flux in powder form, were

used: the latter gave slightly the better results. The flux in each case was an ammonium chloride-zinc chloride mixture. The most suitable nozzle size was found to be that passing 50-60 litres/hr. per mm. thickness of material. A brick backing-plate was used. Photomicrographs and radiographs show that the results obtained were excellent.—H. W. G. H.

**\*Welding of Wires by Condenser Discharge.** O. Zdralek and J. Wrana (*Elektrotech. Z.*, 1935, 56, 579-580).—Wires can be welded by a non-oscillating condenser discharge. Welding takes place in 2 steps, first the ends of the wires fuse to small globules and these then coalesce, producing a larger globule at the joint. The electrical conditions for welding copper, iron, aluminium, Constantan, and Nickel in by this method are given.—B. Bl.

**†The Eleventh International Congress [Rome] on Acetylene and Autogenous Welding and Related Industries.** J. Brillie and (Mlle.) L. Bloch-See (*Rev. Met.*, 1935, 32, 106-119).—A *résumé* of papers presented at the Congress, mainly dealing with ferrous materials. Papers by A. Boutté on the unweldability of nickel and by A. B. Kinzel and A. R. Lytle on brazing with the oxy-acetylene torch are reviewed briefly.—H. S.

**An Early History of Oxy-Acetylene Welding and Cutting in the United States.** Orn T. Weirs (*Welding Eng.*, 1935, 20, (6), 22-24).—An interesting account of the introduction, by Eugene Bournonville, of oxy-acetylene welding and cutting into the U.S.A.—H. W. G. H.

**Internal Stresses in Weld Seams.** — Müllenhoff (*Elektroschweißung*, 1935, 6, 103-108).—The advantages of Mathar's method for measuring internal stress (by boring holes and measuring the deformation round them) are pointed out, and a series of investigations on mild steel welds is described. Special attention is directed to the effect of stressing specimens, beyond the proportional limit, longitudinally to the weld. In a gas-welded specimen, stresses along and across the seam of about 38 and 12½ tons/in.<sup>2</sup>, respectively, decreased to about 6½ tons/in.<sup>2</sup>; in a bare-wire arc-welded piece, a stress of about 28½ tons/in.<sup>2</sup> parallel to the seam decreased to about 16 tons/in.<sup>2</sup>; and, in a sample welded with coated electrodes, stresses of about 25 parallel to, and 19 across the seam, were reduced by half, the stress applied to the specimens being 6½ tons/in.<sup>2</sup>.—H. W. G. H.

**Applications of Resistance Welding.** P. R. Dunn (*Elect. Rev.*, 1935, 117, 177-178).—A general survey.—J. C. C.

**Arc Welding in the Chemical Industry.** P. Stricker (*Arcos, Z. Lichtbogen-schweißung*, 1934, 11, 1127-1129).—The use of electric arc welding in the construction of chemical apparatus is discussed, with examples.—S. G.

## XXI.—INDUSTRIAL USES AND APPLICATIONS

(Continued from pp. 401-405.)

**Effect of Vibration on Steel-Cored Aluminium Conductors.** L. L. Eve (*World Power*, 1934, 21, 247-249; and *Electrician*, 1934, 112, 565; discussion, 565-566; *Sci. Abs.*, 1934, [B], 37, 417).—Vibration is most prevalent with wind velocity of about 5 m.p.h. and where the line is unobstructed. The fatigue breaks have occurred mostly on the 37/0-102 s.c.a. conductor and at entrance to suspension clamps (and sometimes to tension clamps). When the temperature decreases below the erecting temperature the proportion of stress taken by the aluminium of the s.c.a. conductor increases considerably. E. advises overstressing the conductor before sagging the line to ensure against the calculated proportion being severely encroached upon at the lower temperatures. He suggests, as a precaution against fatigue breaks, erecting (1) at a safe tension, or (2) at a tension taking full advantage of the strength of the conductor and equipping the line with dampers.—S. G.

**Method of Damping the Vibrations of Overhead Cables.** M. Preiswerk (*Bull. Assoc. Suisse Elect.*, 1934, 25, 252-253).—See *Met. Abs.*, 1934, 1, 637.  
—S. G.

**Lightness in Truck Design.** Austin M. Wolf (*Soc. Automotive Eng. Preprint*, 1934, 19 pp.).—The advantages of using light aluminium in the construction of the chassis of trade motor vans, &c., are discussed with reference to numerous examples. Tables of the mechanical properties and applications of aluminium alloys for trucks and truck bodies are included.  
—A. R. P.

**Aluminium Castings and Their Uses in Modern Vehicle Construction.** Oskar Summa (*Automobiltech. Z.*, 1935, 38, (12), 297-301).—A tabulated comparison of important physical and mechanical properties of aluminium with those of magnesium, copper, tin, iron, and steel, is followed by a summarized account of aluminium alloys containing silicon, magnesium, copper, and/or zinc. Their properties in the cast condition are tabulated. Their freedom from thermal distortion makes them especially suitable for use in motor engines. Defects resulting from faulty preparation are described and classified, with special emphasis on the technique of remelting. Compositions and properties of Nural, 142, copper-Silumin, and Silumin-Beta are given, and the thermal changes of the last-named are described.—P. M. C. R.

**Cast Aluminium, a German Material.** W. Hartl (*Automobiltech. Z.*, 1935, 38, (12), 301-307).—An account is given of the physical and mechanical properties of certain light alloys—Pantal, Silumin, Copper-Silumin, and Silumin-gamma—all of which are said to be especially suited for use in the cast condition. Cleaning, finishing, and protection methods are described, and data are given regarding the corrosion of Silumin and of pure aluminium in liquid fuels, including some proprietary mixtures. Detailed instructions are given for the substitution of light alloy for copper and nickel parts.  
—P. M. C. R.

**Twenty-Five Years of Light Metal Pistons.** E. Mahle (*Aluminium*, 1935, 17, 355-369).—A review of the development of the design of light metal alloy pistons during the past 25 years.—A. R. P.

**\*Aircscrew Oscillations.** M. Hansen and G. Mesmer (*Z. Flug. u. Motor.*, 1933, (11); and (translation) *Aircraft Eng.*, 1935, 7, 65-69).—Theoretical considerations on natural frequency and torsional oscillations are supplemented by experimental data on models. The results suggest that there is considerable danger of serious resonance effects between overtone oscillations due to blade bending and the pulsations of the engine particularly with thin metal aircscrews having low natural frequencies.—H. S.

**Aluminium in Shipbuilding.** — Sterner-Rainer (*Aluminium*, 1935, 17, 370-374).—The advantages of magnesium-aluminium alloys in shipbuilding are discussed, and the technique of melting, casting, and working them is described. Surface protection of aluminium alloys by plating with pure aluminium and by coating them with oxide by the Eloxal process is briefly considered.—A. R. P.

**\*Formaldehyde in Canned Marine Products [in Aluminium and Tinplate Cans].** Gulbrand Lunde and Erling Mathiesen (*Indust. and Eng. Chem.*, 1934, 26, 974-976; and (abstract) *Light Metals Rev.*, 1934, 1, (12), 242).—Formaldehyde can be detected in the distillate from fresh or canned marine products, and tests show that there is no difference between the amounts formed in tinplate or in aluminium vessels after various periods of storage. The formation of formaldehyde, which was thought to be caused by reaction between the fish and the containers, does not increase during long storage.  
—R. B. D.

**Recent Developments in Metallurgy and Design of [Bronze] High-Speed Motor-Boat Propellers.** T. F. W. Meyer (*Soc. Automotive Eng. Preprint*, 1933, 17 pp.).—A description is given of the development, construction, and per-

formance of Federalite bronze propellers used in the American competitor for the motor-boat trophy.—A. R. P.

**The Use of Lead in the Oil Industry.** G. O. Hiers (*Amer. Metal Market*, 1935, 42, (133), 3, 5).—Lead-lined tanks and pipes are used widely in the oil industry in refining processes involving the use of sulphuric acid. The properties of chemical lead, antimonial lead, and tellurium-lead are reviewed and the suitability of these 3 grades of lead for various purposes in the oil refinery is indicated.—A. R. P.

**\*Stability of Lead Alloy Anodes in the Electrolytic Production of Zinc.** P. S. Titov and I. N. Nikonov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (6), 53-62; *C. Abs.*, 1935, 29, 4269).—[In Russian.] A study was made of the stability of anodes made of binary lead alloys containing various amounts of bismuth, antimony, arsenic, and silver, in the electrolysis of zinc, in ordinary conditions of operation, and the results were compared with those obtained from experiments with pure lead anodes. Lead-bismuth anodes containing 0.5-10% bismuth were less stable than pure lead anodes. Alloys containing 0.5-10% antimony were somewhat more stable than pure lead at 400 and less stable at 1000 and 1200 amp./m.<sup>2</sup>. The amount of arsenic added to the lead varied from 0.5 to 3.0%. At 400 amp./m.<sup>2</sup> the stability increased for the 0.5 and 1.0% arsenic alloys, decreased for the 2.0% and increased again for the 3.0% alloy. At 1000 and 1200 amp./m.<sup>2</sup> the stability of all the arsenic alloys was less than that for pure lead. Addition of silver, 0.5-3.0%, to the lead anodes increased the stability considerably (cf. Ageenkov and Sosunov, following abstract). The stability curve reaches a maximum at 2.5% silver, where the anodes last 20-25 times as long as in the case of pure lead. Small additions of antimony up to 0.5%, of arsenic up to 0.5%, of bismuth up to 0.2%, or of bismuth and arsenic together, 0.1 and 0.4%, respectively, produced no deleterious effect on the lead-silver anodes. The presence in the sulphuric acid bath of zinc sulphate, manganese sulphate and glue had no appreciable effect on the stability of the anodes.—S. G.

**\*Effect of Anode Composition on Zinc Electrolysis.** V. G. Ageenkov and S. L. Sosunov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (5), 61-74; *C. Abs.*, 1935, 29, 4269).—[In Russian.] Cf. preceding abstract. Lead anodes containing small amounts of silver or calcium, or both, were compared with pure lead anodes in the electrolysis of zinc ore from a sulphuric acid bath. The composition of the lead used was: arsenic, trace; antimony, 0.00107-0.00910; bismuth, 0-0.002; copper, trace-0.00130; iron, 0.00192-0.00143; lead, 99.994-99.996%. The amount of silver in the silver-lead anodes or silver-lead-calcium anodes varied from 0.462 to 1.987% and of calcium, from 0.28 to 1.06%. The electrolyte contained on the average 60 gm. zinc, 100 gm. sulphuric acid, 0.174-0.36 gm. manganese, and 0.007 gm. iron per litre; the temperature varied from 21.0° to 29.8° C., voltage 3.25-3.75, and anode density 400 amp./m.<sup>2</sup>. The presence of silver improves the chemical resistance of the anode, lowers the voltage, and reduces the amount of lead in the zinc deposit. The presence of calcium in addition to silver intensifies the effects due to silver. The mechanical strength of the lead anode is increased by the addition of silver and particularly calcium.—S. G.

**Plumbers' Metal.** — (*Mech. World*, 1935, 97, 177).—Notes prepared and communicated by British Insulated Cables, Ltd., for the information and assistance of those using plumbers' metal (tin 33.3, lead 66.7%)—F. J.

**[Molybdenum] Crucibles for High-Temperature Work.** — (*Mech. World*, 1935, 97, 237, 286, 383-384, 433, 518, 569).—The pros and cons of the application of molybdenum to large-scale work in the form of crucibles are discussed by correspondence.—F. J.

**Properties of Nickel and Nickel Alloys in Caustic Processing Equipment.** Takashi Okamoto (*Japan Nickel Information Bureau, Tech. Information B-19*, 1935, 50 pp.).—[In English and Japanese.] From a paper read before

the Society of Chemical Industry, Japan. The behaviour of materials in dilute and concentrated caustic solutions and in fused caustic soda is reviewed, and the advantages of nickel and its alloys are pointed out. The selection of the most suitable materials for various parts of caustic processing plant is discussed, and the properties of those recommended are given in detail.

—H. W. G. H.

**Nickel Alloys in the Paper-Making Industry.** A. B. Everest (*World's Paper Trade Rev.*, 1935, 103, 831-836, 876, 878, 996-1000, 1034, 1036).—The applications of nickel alloys in the paper-making industry are described.

—S. G.

**Nickel and Nickel Alloys in the Refining of Crude Oil.** R. W. Müller (*Oil u. Kohle*, 1935, 11, 504-507).—Alloys containing a high proportion of nickel, especially Monel metal, are much used in chlorination plant and in coolers and condensers where sea-water is used. Monel metal is used for bolts, piston-rods, tubing, sheet and valve and pump parts. The 70-30 copper-nickel alloy is employed in condensing and heating plant, as is Inconel, which combines high resistance to corrosion with unusual freedom from surface carbonization. Some steel parts are nickel-plated as a protection against attack by acids, alkalis, or petrol.—P. M. C. R.

**Alkaline Accumulators.** Guy Malgorn (*Électricien*, 1934, 45, 126-132).—A description of the manufacture, characteristics, and applications of nickel-iron and nickel-cadmium accumulators, and a discussion of their advantages and disadvantages as compared with lead accumulators.—C. E. H.

**Correlation of Viscosity Measurements with Flow of Glass.** A.—Development of the Platinum Alloy-Lined Die: the Die of Unvarying Diameter. H. K. Richardson (*J. Amer. Ceram. Soc.*, 1934, 17, 236-239).—S. G.

**Silver for Chemical Equipment.** C. H. S. Tupholme (*Chem. Industries*, 1935, 36, 527-528).—The action of various chemicals on silver is described and the use of silver in chemical plant for handling foodstuffs, acetic acid, and hydrofluoric acid is briefly discussed.—A. R. P.

**Bearing Metals.** F. K. von Göler and G. Sachs (*Metalgesellschaft Periodic Review*, 1935, (10), 3-10).—[In English.] The structure and most important physical and mechanical properties of tin-, lead-, and cadmium-base bearing metals, of lead-base alloys containing alkali and alkaline earths, of lead bronzes and of zinc- and aluminium-base bearing metals are reviewed, and the selection of suitable alloys for specific purposes is discussed.—v. G.

**Engine Bearings from the Designer's View-Point.** Alex Taub (*J. Soc. Automotive Eng.*, 1934, 31, 385).—Curves are given and discussed showing the relation between loading, shaft speed, and life of various bearing metals.—A. R. P.

**The Bearing Manufacturer's View-Point on Engine Bearings.** D. E. Anderson (*Soc. Automotive Eng. Preprint*, 1932, 15 pp.).—The max. bearing pressure and oil temperature permissible in bearings are determined by the properties of the Babbitt metal and the lubricating oil. The best grade of Babbitt has a compressive resistance of about 2000 lb./in.<sup>2</sup> at 175° C. and a bearing having a load of 1000 lb./in.<sup>2</sup> produces a hydrostatic pressure of nearly 2000 lb./in.<sup>2</sup> at its centre; since tests have shown that the metal temperature in the bearing is about 55° C. above that of the oil supplied to it the Babbitt temperature will be within the range at which the metal becomes plastic and fails when the oil temperature is 120° C. The P-V factor of the bearing must, therefore, be such that the mean unit pressure is less than 1000 lb./in.<sup>2</sup> and the oil temperature must be below 120° C. Thus at 4000 r.p.m. the P-V factor must not exceed 40,000. At higher engine speeds the bearings must be made of lead-copper or silver alloys to withstand the pressure and heat.—A. R. P.

**Modern Alloys in Marine Engineering and Shipbuilding.** J. W. Donaldson (*Metal Treatment*, 1935, 1, 59-62).—A general review. The non-ferrous

alloys mentioned include modified aluminium-silicon alloys for hull work; Monel metal for interior construction, linings for trawler fish-holds, and valves; 70:30 copper-nickel alloys and aluminium-brass for condenser tubes; and high-duty copper alloys.—J. C. C.

**Motor Manufacture in the Light of Available Unrestricted Materials.** Erich Gründger (*Automobiltech. Z.*, 1935, **38**, (12), 294-298).—The use for certain industrial purposes of copper, nickel, lead, and tin, as well as of chromium, nickel, or cobalt plating, is now forbidden in Germany. A list is given of important automobile parts affected by these restrictions, and alternative possibilities are considered. Where possible, ferrous materials should be avoided in view of their liability to corrosion, though zinc coatings are recommended for some purposes. Where synthetic resins are unsuitable, there is a wide field of light metals and alloys, and a summary is given of the available range of mechanical properties, of anti-corrosive measures, and of special means of protecting magnesium alloys.—P. M. C. R.

**The Piping Installation.** A. V. Knowles (*Motor Boat*, 1935, **63**, 125).—Materials and lay-out for the water-circulating system of a motor-boat are discussed. Copper is considered to be the best material for all water, fuel, and oil pipes, and where some filling is necessary with such pipes lead is recommended. Flanges and unions on copper pipes should be of brass and should be brazed or hard (silver) soldered on. Soft soldering is not recommended for marine work.—J. W. D.

## XXII.—MISCELLANEOUS

(Continued from pp. 405-406.)

**The Science of Metallurgy in Technology and Research.** Paul Röntgen (*Aachener Akad. Reden I*, 1934, 16 pp.).—A review of recent developments in non-ferrous metallurgy, with especial reference to the production of electrolytic zinc, aluminium, and beryllium, and to progress in the heat-treatment of non-ferrous metals.—A. R. P.

**State of Non-Ferrous Metallurgy in Foreign Countries.** V. Silinskaya (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (4), 92-100).—[In Russian.]—S. G.

**Cadmium and Its Outlook in the U.S.S.R.** N. M. Siluyanov (*Zvetnye Metally (Non-Ferrous Metals)*, 1934, (10), 148-154).—[In Russian.]—S. G.

**The Order Book of the New Wire Company, Cheadle, 1788-1831.** L. St. L. Pendred (*Newcomen Soc. Advance Copy*, 1935, Mar., 18 pp.).—The activities of an old English company making copper and brass wire are surveyed between 1788 and 1831 by extracts from the order and minute books.—A. R. P.

**The London Lead Company, 1692-1905.** A. Raistrick (*Newcomen Soc. Advance Copy*, 1934, Apr., 38 pp.).—The development and activities of the company in the production of lead and silver from mines in the north of England are reviewed, and a brief account is given of the contribution of the company to the technique of mining and smelting lead ores.—A. R. P.

**Epidemiology of Lead Poisoning.** A. J. Lanza (*J. Amer. Med. Assoc.*, 1935, **104**, 85-87; *C. Abs.*, 1935, **29**, 4852).—A daily dose of from 1.5 to 2 mg. is dangerous. Lead in the form of vapour is more dangerous than as dust.—S. G.

**Control of Lead Poisoning in the Worker.** Elston L. Belknap (*J. Amer. Med. Assoc.*, 1935, **104**, 205-211; *C. Abs.*, 1935, **29**, 4852).—A successful working technique is outlined for the medical control of poisoning in industrial occupations.—S. G.

**Manganese: Its Minerals and Industry.** Maurice Dérivé (*Bull. anc. Écoles École Bréquet*, 1933, (86), 1-20).—A general review.—S. G.