2561 to Dr.

### PHYSICS ABSTRACTS

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539.32

2402

**EFORMATION** 

Measurement of the modulus of rigidity of solid and liquid bodies. KORNFELD, M. J. Phys., USSR, 8, 1, pp. 4-8, 1944.—Apparatus is described for measuring the rigidity modulus of crystalline and amorphous bodies, and the method may be applied without limitation to substances whose viscosity exceeds 106 poises. The theory of the method is given. L. S. G. 539.32:532.13:666.1

The determination of the elastic and viscous properties of glass at temperatures below the annealing range. JONES, G. O. J. Soc. Glass Technol., 28, pp. 432-462, Dec., 1944.-A bending method was used for observing the development of strain under applied stress. Down to quite low temperatures the whole of the strain is made up of a purely viscous part, plus a completely reversible elastic part. The reversible strain may be divided into an instantaneous component and a delayed component which appears rapidly at first and then approaches asymptotically to a max. A method was developed for the analysis of the strain/time curves, based on the observation that only that part of the delayed strain which had already appeared under load was later recovered on removal of the load, and on the assumption that the form of the decay function was similar for both deflection and recovery. The total delayed elastic strain for a given applied stress is a function of the temperature, increasing from 3% of the instantaneous elastic strain at 200° to 75% at 444° ( $T_g$  was 534°). Above this temperature it became increasingly difficult to separate the instantaneous and delayed parts of the elastic strain. The true viscosity at 350° was 1017.8 poises.

539.32: 534.321.9: 548.0 see Abstr. 2565

539.32: 548.0 see Abstr. 2564

539.32 : 553.621 2404

Elastic constants of quartz. RAO, B. R. Curr. Sci., 14, p. 17, Jan., 1945.

539.378.3 : 548.526 : 669.14 240

Solution of the diffusion equation applicable to the edgewise growth of pearlite. BRANDT, W. H. J. Appl. Phys., 16, pp. 139-146, March, 1945.—The diffusion equation is transformed to a set of co-ordinates moving with the pearlite interface and a solution applicable to the problem obtained in the form of an infinite series of terms. Using the first 3 terms, the edgewise velocity of pearlite growth is calculated for a plain carbon eutectoid steel using data most of which are obtained by extrapolation. The values obtained agree with values for the rate of pearlite nodule growth determined by Hull, Colton, and Mehl. The velocity increases with decreasing temperature, as expected, and it is shown that this is caused by the change in the solubilities of ferrite and cementite in austenite with temperature. The theory predicts curved ferriteaustenite and cementite-austenite interfaces and the carbon concentration in austenite is shown to vary across each of these interfaces.

539.379: 539.501 2406

Rise of temperature on fast stretching of Butyl rubber. Dart, S. L., and Guth, E. J. Chem. Phys.,

13, pp. 28-36, Jan., 1945.—[See Abstr. 1509 (1943)]. Study of the rise in temperature on fast stretching of Butyl gum and tread stocks throws light on crystallization, internal friction, and carbon-black reinforcement as functions of elongation and other variables. For low clongations, the low-heat curves show thermo-elastic inversion. They also show that internal friction is much larger for Butyl than for Hevea. For high elongations, the high-heat curves enable one to follow the late and abrupt crystallization, characteristic for Butyl stocks. The carbon-black reinforcement of Butyl is followed for the whole range of elongations up to the breaking point. Specific carbon-black reinforcement takes place in the range of elongations before crystallization of the gum stock, but the shortening of the breaking elongation by the black excludes to a large extent the added reinforcement through crystallization.

539.381 2407

Wave equation for finite elastic strains. NADAI, A. J. Appl. Phys., 16, p. 313, May, 1945.—[See Abstr. 2620, 2622 (1944)].

539.382:620.172.22.05

2408

The yielding phenomenon of metals. IV-VI. Welter, G. Metallurgia, Manchr, pp. 309-315, April; 31-36, May, and pp. 80-84, June, 1945.—[Abstr. 1817 B (1945)].

539.384.4:674.03

2409

Hardwood columns subjected to loads of short duration. BULL, M. G. J. Instn Engrs Aust., 17, pp. 50-58, March, 1945.—A discussion of theory as generally applied to the testing of timber is followed by an analysis of tests on columns of Australian hardwoods. The max. load-carrying capacity may be found by assuming the limiting physical properties of the column in conjunction with a relationship between direct and bending stresses at max. load as found by Warren for spruce. It is concluded that the mean behaviour of hardwood columns under combined bending and compressive stresses is substantially similar to that of softwood columns. G. E. A.

Bending of curved thin tubes. Beskin, L. J. Appl. Mech., 12, A1-A6, March, 1945.—The stress distribution in curved tubes, and its influence on the rigidity,

are examined in the case when the ratio of the radius of the centre line to the radius of the tube is great. When this ratio is small, the results relative to rigidity remain fairly accurate, while the results relative to stress distribution are incorrect and require a more

complete analysis.

539,385.08: 679.5: 621.383.4

A rapid photo-electric optical distortion tester for plastic windows. Sowerby, J. McG., and Walton, W. H. J. Sci. Instrum., 22, pp. 71-74, April, 1945.— Describes an instrument suitable for use in the factory or industrial test room for the "pass-reject" testing of the optical distortion of small moulded plastic windows. The instrument incorporates a "Schlieren" optical system by which faulty windows deflect light off a stop on to a photocell so that the photocell current is a measure of the optical distortion. By

means of an electronic circuit incorporating a hard valve trigger (self-resetting), the photocell current is made to operate an indicator lamp. Standardization is effected by adjustment of the circuit just to reject a glass distorting standard. The instrument derives its supply from the a.c. mains and sufficient stabilization is incorporated so that only infrequent standardization is needed. A testing rate of 24 windows per min. is easily attained.

539.388.1 : 548.73 see Abstr. 2591

539.388.8:678 2412

Swelling of synthetic rubbers in mineral oils. Powers, P. O., AND BILLMEYER, B. R. Industr. Engng Chem., 37, pp. 64-67, Jan., 1945.—The aniline point of hydrocarbon oils and solvents is a satisfactory index of the swelling of oil-resistant synthetic rubber compositions. The log. of the % volume increase varies inversely with the aniline point up to 100% swelling. The slope of the swelling curve is characteristic of each synthetic rubber and is not affected by loading, temperature or degree of cure. Slight swelling does not greatly decrease the tensile strength, but above 100% swelling the strength is greatly reduced.

539,388.8: 678.7

Identification of raw and vulcanized rubber-like polymers. II. Determination of swelling ratios. Parker, L. F. C. J. Soc. Chem. Ind., 64, pp. 65-67, March, 1945.—The ratios of the swellings of vulcanized rubber-like polymers in benzene, light petroleum (b.p. 40-60°), and aniline, at 25°C., are different for different polymers, but within a well-defined narrow range for each polymer, the ratios being independent of the state of vulcanization and the nature and amount of the fillers. Hence the ratios for an unknown sample may be utilized in its identification, preferably by plotting the values on a graph showing the ratios for the common types of rubber-like polymers.

539.389 2414

A theory of the Calender effect in rubber-like materials. BILMES, L. Trans. Faraday Soc., 41, pp. 81-87, Feb., 1945.—The facts relating to the Calender effect are briefly stated and illustrated by experimental data obtained with calendered plasticized polyvinyl chloride sheet. The nature of rubber-like deformation is outlined. A theory of the Calender effect is proposed in terms of a mechanical model whose relation to molecular structure is indicated. The theory postulates that temperature-dependent yield values be associated with the mechanisms of high elastic deformation and plastic flow.

539.389.3 : 539.501 2415

Pseudo-plastic phenomena with vulcanized rubber. WILDSCHUT, A. J. Physica, 's Grav., 10, pp. 571-584, July, 1943.—The variations of plastic flow and relaxation with time are studied. As a result, the rate of flow (v) is defined as the ratio of the flow at const. tension to the log. of the time, and the rate of relaxation as the ratio of the decline in tension at const. elongation to the log. of the time. The rates thus defined are independent of the time. The rate of flow decreases rapidly with rise in temperature, and this is explained by considering the influence of time on crystallization and of crystallization on elastic tension. When v is measured at increasing values of the tension  $(\sigma)$  and v is plotted against  $\sigma$ , the result is a

curve with a strong max. This is explained by the crystallization of the rubber. A study of such curves shows that, at room temperature, a large part of the total flow is reversible and must be considered as pseudo-flow, which is caused by a retardation in the crystallization and diminishes at increasing temperature. A consequence is that the total rate of flow passes through a min. as the temperature increases. The phenomenon of pseudo-relaxation is also studied.

L. S. G. 2416

Effect of temperature on strength of laminates. NORELLI, P., AND GARD, W. H. *Industr. Engng Chem.*, 37, pp. 580-585, *June*, 1945.—[Abstr. 1969 B (1945)]. 539.4.01: 679.5 2417

Effect of some environmental conditions on the mechanical properties of cellulose acetate and cellulose nitrate plastic sheets. Lawton, T. S., Jr., Carswell, T. S., and Nason, H. K. Trans. Amer. Soc. Mech. Engrs, 67, pp. 23-30, Jan., 1945.—The effect of ambient temperature, relative humidity, testing speed, and sheet thickness on the tensile properties, and of temperature on the compressive, flexural, shear, and impact properties of cellulose acetate and cellulose nitrate plastic sheets, is shown in detail and the general nature of these effects is summarized.

539.4.016.3:669.721 2418

An exploration of the problem of superheating in magnesium-base alloys. Fox, F. A., and Lardner, E. J. Inst. Met., 71, pp. 1-22, Jan., 1945.—The grainrefining effect is confined to alloys containing Al, and tendencies to grain-coarsening are introduced if the superheating time is too long or the temperature too high. Data are given relating tensile properties to grain-size in Al-containing alloys. The microstructures of superheated alloys differ characteristically from those of the unsuperheated material, and this difference persists even after solution treatment, the unsuperheated alloy giving a mixed grain-size. Grain-sizes are recorded for an alloy, superheated and unsuperheated, cast as rods of various cross-sections. Stirring just before casting does not eliminate the grain-refinement effect of the superheating. 539.415.2:621.93.01 2419

Mechanics of the metal cutting process. II. Plasticity conditions in orthogonal cutting. MERCHANT, M. E. J. Appl. Phys., 16, pp. 318-324, June, 1945.—[Abstr. 2295 B (1945)].

539.42: 541.64 see Abstr. 2524

539.42: 677.2: 539.216.1 see Abstr. 2385

539.42: 677.2: 539,216.1 see Abstr. 2383 539.43: 669,018

Dependence on stress of damping capacity of alloys. GEMANT, A. Mech. Engng, N.Y., 67, pp. 33-38, Jan., 1945.—A consideration of damping as caused by the alternating plastic flow that takes place during vibration. The decrement is converted into transient flow resistance which is plotted against the stress for a number of alloys of steel, Cu and Pb. Certain of these have damping characteristics with pronounced dependence on the stress, and this characteristic is particularly promoted by additions of Si and Ni. Heat treatment is not considered.

G. E. A.

On fatigue failure under triaxial static and fluctuating stresses and a statistical explanation of size effect. FOWLER, F. H., JR. Trans. Amer. Soc. Mech. Engrs,

67, pp. 213-216, April, 1945.-A criterion of safe fatigue characteristics of manufactured units is proposed. A basic theory of failure is recommended which is a modification of Drowan's [Abstr. 1906 (1939)]. Explanations of the effect of specimen size and the stress gradient on the endurance limit may be useful in avoidance of the possibility of failure or in evaluating the probability that a failure will occur.

G. E. A.

539.431:678

2422 Fatigue failure of GR-S tread stocks. WINN, H., AND SHELTON, J. R. Industr. Engug Chem., 37, pp. 67-70, Jan., 1945.—The effect of the state of cure, temperature of test and oxidation, on the fatigue failure in rubber during flexing in air and in N<sub>2</sub> containing small amounts of O2 is investigated experimentally.

539.434: 679.5

Creep properties of moulded phenolic plastics at elevated temperatures. GAILUS, W. J., AND TELFAIR, D. Trans. Amer. Soc. Mech. Engrs, 67, pp. 253-258, May, 1945.—Tension creep data are reported up to 1 000 hr. and recovery data up to 250 hr. Modulusof-elasticity values were obtained at the beginning and end of the 1 000 hr. tests. Studies included the effect of moisture content on creep properties.

539 501 2424

Experiments on plasticity. I-II. MACEY, H. H. Trans. Brit. Ceram. Soc., 43, pp. 5-28, Jan., 1944.-Two simple methods of experiment on plastic clay are given and the data obtained by these methods discussed. Clay bars are subjected to constant tensional and compressional stresses less than those necessary to produce continuous flow. It is shown that clay is not elastic, but deforms plastically even at the smallest stresses. De-aired clay bars are extended at a constant speed and a continuous record of the stress involved is obtained. A wide range of speeds is used and it is shown how the behaviour of the bar varies with the speed. The extension to max. stress increases as the rate of extension is increased. Considerable variations in behaviour of apparently identical bars are found, but the stress/extension curves agree with the formula  $f = B(1 - e^{-aE})$ , where f is the true stress per unit area, E is the extension and a and B are constants for any one experiment.

2425

Systems manifesting superposed elastic and viscous behaviour. Tobolsky, A. V., and Andrews, R. D. J. Chem. Phys., 13, pp. 3-27, Jan., 1945.-A general molecular theory describing the behaviour of matter under stress is discussed and related to previous investigations and to experimental observations. Experimental results on relaxation and creep of rubbers are interpreted in terms of modern structural concepts. These substances exhibit 3 regions of stress/temperature/time dependence. At intermediate temperatures there exists a region of relative stability in which the statistical-thermodynamic theory of rubber elasticity is valid. At elevated temperatures relaxation and creep are caused by chemical changes involving the rupture and formation of primary valence bonds. These chemical changes, which are responsible for the ageing of rubber, can be isolated

and studied by appropriate experimental techniques. At low temperatures relaxation and creep are caused by the slipping of secondary interchain bonds which are breaking and reforming in times comparable with experimental times of measurement.

539.501 2426

Significance of power-law relations in rheology. CAFFYN, J. E., AND BLAIR, G. W. SCOTT. Nature, Lond., 155, pp. 171-172, Feb. 10, 1945.-[See Abstr. 2427 (1945)].

539.501

Power laws for the dependence of strain on time and WHITEHEAD, S. Rep. Brit. Elect. Allied Industr. Res. Ass., Ref. L/T151, 10 pp., 1945.-A Laplace transform method is used to show that it is impossible for a linear system to give rise to a power law; but such a law may approximate the behaviour over a limited range of time. If we abandon the hypothesis of linearity but assume the phenomena are still superposable we find that the law of variation with time is given by an integral equation. Since the latter is the limit of a system of linear equations it follows from the previous work that a power law cannot apply to superposable phenomena. There is a section on the mechanism of relaxation, and the report has particular reference to the rheological properties of dielectrics [See Abstr. 2426 (1945)].

L. S. G.

539.501 2428

A classification of rheological properties. Reiner, M. J. Sci. Instrum., 22, pp. 127-129, July, 1945.-An outline is given of the method of describing the behaviour of complex materials under strain in terms of dashpots, springs, etc., various simple combinations ascribed to St. Venant, Kelvin and others being described. A proposal is made to classify rheological properties in terms of the ideal bodies possessing them. The method of application to a new type of material is illustrated by considering flour dough, as examined by Scott-Blair and Schofield. Two rheological properties, dilatancy and strength, are excluded from the proposed scheme.

539.501: 532.133 see Abstr. 2173 539.501: 539.379 see Abstr. 2406

539.501: 539.389.3 see Abstr. 2415

539.501:662.66 2429

Plastic characteristics of coal. Brewer, R. E. Industr. Engng Chem., 36, pp. 1165-1168, Dec., 1944.—A detailed study of chemical and petrographic composition and physical properties of 19 low-, medium-, and high-volatile A bituminous coals in relation to Gieseler max. fluidity. Factors of chemical and petrographic composition found to contribute toward increasing the fluidity of a coal above that expected from its rank are high anthraxylon and translucent attritus contents and the presence of cannel coal. For coals of similar rank, those of higher physical strength generally show a higher max. fluidity. Factors that reduce the fluidity of a coal below that expected from its rank are high contents of oxygen, ash, opaque attritus, and fusain.

539.501:678 2430

Plastication and processing of GR-S. VILA, G. R. Industr. Engng Chem., 36, pp. 1113-1119, Dec.,

1944.—Viscosity or plasticity measurements bear a direct relation to the power required and the heat generated while tubing GR-S, but do not correlate with tubing rates. The latter depend upon worm speed and the surface condition of the mixed compound, smoother surfaces tending to give faster speeds. The increase in cross-sectional area upon tubing bears a direct relation to surface smoothness, smoother compounds exhibiting less swell. possible to predict tubing characteristics from the plasticity or viscosity of the finished compound in conjunction with an estimate of its relative surface smoothness. The latter quality may be achieved by hot breakdown, increasing softener ratios, or increasing pigment ratios. High-temperature plastication, however, exerts a deleterious effect on the physical properties of the vulcanizate. There is evidence that a higher degree of benzene solubility in the raw polymer may result in a more favourable processing history, but additional examples are required before conclusions are justified. working with normal GR-S, cold milling is found to increase benzene solubility whereas hot milling or Gordon plastication reduces it and, presumably, results in a gelled polymer.

539.536 2431

Abrasion of porcelain balls. Chesters, J. H. Nature, Lond., 155, p. 485, April 21, 1945.

539.536: 676.3 2432 Determination of the erasing qualities of paper. CODWISE, P. W. Paper Tr. J., 120, TAPPI Sect., pp. 73-76, Feb. 22, 1945.—The (Howard Smith) mechanical erasure tester described consists of a uniformly moving and controlled abrading surface (one of a series of standardized grit abrasive cloths), which is made to rub the paper surface under a total head wt. of 327 g., at 120 strokes/min., by means of a motor-driven flywheel with an eccentric arm. The test is made on a cross-hatched line on the paper drawn with a drawing pen set to 0.0001 in., and using the TAPPI standard (T431 m-41) ink. Results are tabulated for each side of 5 varied papers. In assessing the erasing quality of paper, the time to achieve erasure, the smoothness and ink-resisting properties of the surface after erasure, the amount of paper removed by the crasure, and the tendency of the paper to "fuzz up" during erasure, should all be taken into account. J. G.

539.56 : 548.0 2433

A statistical theory of the brittle strength of real crystals. Frenkel, J. I., and Kontorova, T. A. J. Phys., USSR, 7, 3, pp. 108-114, 1943.—A theory is developed which is more strict and accurate than previous theories. The probability of encountering a given value for the brittle strength of a crystalline material is calculated and from this the most probable value of the brittle strength is deduced, this depending upon the volume of the specimen. The results are compared with those of Weibull [Proc. Roy. Swedish Inst. Engng Res., No. 151, 1939].

L. S. G.

539.56: 666.1

The mechanism of brittle rupture in glass. Mur-GATROYD, J. B. J. Soc. Glass Technol., 28, pp. 406-431 (T), Dec., 1944.—The suggestion is made that Griffith "flaws" consist of pockets of quasi-viscous material surrounded by larger 3-dimensional aggregates, and that, when the pocket material has yielded to an applied stress, the fact that it then carries no load makes the pocket the equivalent of a "hole" in the glass. The time elapsing before breakage occurs is explained as being due to the time required for the stress in the quasi-viscous material to relax. By means of a model representing the stress concentration at a pocket, an equation is derived relating the breaking strength with time.

539.574:667.7 2435

Nitrocellulose lacquers. Koch, W., Phillips, H. C., AND Wint, R. Industr. Engng Chem., 37, pp. 82-86, Jan., 1945.—Non-volatile content can be increased in nitrocellulose lacquers by using lower-viscosity nitrocelluloses and larger proportions of non-oxidizing alkyd resins with a more active solvent, or by applying the lacquers at temps. up to 70°C. The gain in non-volatile content contributed by each factor is cumulative

539.61 2436

Adhesion of lubricated metals. Bowden, F. P., and MOORE, A. J. W. Nature, Lond., 155, pp. 451-452, April 14, 1945.—An examination of the contours of solid surfaces shows that the real area of contact between two surfaces is only a small fraction of the actual apparent area of contact, so that quite small loads develop large pressures at the few points of contact and local adhesion and welding occurs at these points. A curved Cu slider loaded by 4 kg was moved over a mild steel surface at 0.01 cm. sec.-1, and, when the surface was clean, many tiny particles of Cu were left adhering to the steel, the Cu being firmly welded to the steel. The shearing away of this Cu caused a plucking up of the steel surface, with a consequent wearing away of the steel, and the coeff. of friction was 0.6-0.7. When the surface was lubricated, the friction dropped to 0.15, but the adhesion of the Cu to the steel surface, although much less than in the dry test, was still quite considerable. When Cu and Pt were used, the surface density of the Cu adhering to dry Pt was 2 × 10<sup>-5</sup> gm./mm.<sup>2</sup> of track, while with lubricated Pt the value was  $1.7 \times 10^{-7}$  gm./mm.<sup>2</sup> A. C. W.

539.62: 532.516 2437

The Hardy coefficient for instrument oil. BLOOMEN-THAL, S. Proc. Amer. Phys. Soc., Chicago, Dec. 1 and 2, 1944. Abstr. in Phys. Rev., 67, p. 61, Jan. 1 and 15, 1945.—The coefficient of static friction  $\mu$  for glass on glass was measured by the inclination method on an apparatus which functions automatically. A number of instrument oils, suggested as possible substitutes for genuine porpoise jaw oil, were tried. Addition of 1% glyceryl monoricinoleate to a pure lubricating fluid caused a decrease in  $\mu$  from 0·26 to 0·13 because of boundary layer formation;  $\mu$  for M 56 A Elgin Watch Oil was 0·17 when 100 g. total load is used.

Oil was 0.17 when 100 g, total load is used. 539.62: 669.14

Metallurgical transformations in metal surfaces under conditions of boundary lubrication. SAKMANN, B. W. Proc. Amer. Phys. Soc., New York, Jan., 1945. Abstr. in Phys. Rev., 67, p. 200, March 1 and 15, 1945.—The surfaces of steel flats run-in on rotating cast-iron discs under lubricated conditions may show metallurgical transformations. The transformed

2438

material is considerably harder than the underlying steel and protrudes over initially smooth surfaces by several microns. The experiments include etching tests, microhardness tests, X-ray diffraction pictures, different heat treatments of the transformations, and inspection of the micro-structure of the transformed material and the transition region between the transformation and the bulk material of the steel. From the value of the friction force on the stationary flat and the linear velocity of the rotating disc, it is possible to determine the heat produced during the running-in process.

541.113 2439

High-temperature heat contents of sodium metasilicate and sodium disilicate. NAYLOR, B. F. J. Amer. Chem. Soc., 67, pp. 466-467, March, 1945.-Heat contents of Na<sub>2</sub>SiO<sub>3</sub> and Na<sub>2</sub>Si<sub>2</sub>O<sub>5</sub> were determined between 298·16° and 1 773°K. The heat of fusion of Na<sub>2</sub>SiO<sub>3</sub> was determined, but glass formation precluded a value being obtained for Na<sub>2</sub>Si<sub>2</sub>O<sub>5</sub>. Algebraic equations summarizing the results are given. W. R. A.

541.122.2:669.215 2440

The solubility of certain metals in gold. OWEN, E. A., AND ROBERTS, E. A. O'D. J. Inst. Met., 71, pp. 213-254, May, 1945.—The α-boundaries of 12 gold alloy systems (Be, Al, Zn, Ga, Ge, As, Cd, In, Sn, Sb, Hg, Pb) were investigated by the X-ray method. Parameter/composition curves are drawn for the α-phase alloys of each system in which the solute element has a range of solubility. The effect of various factors on maximum solubility is briefly discussed. [See Abstr. 1514 (1939)].

541,123,1,012,6 Vapour-liquid equilibrium constants for benzene, toluene, and methylcyclohexane. DRICKAMER, H. G., AND BRADFORD, J. R. Industr. Engng Chem., 36,

pp. 1144-1145, Dec., 1944. 541.123.2

Distribution equation. CAMPBELL, J. A. Industr. Engng Chem., 36, pp. 1158-1161, Dec., 1944.—The application of the Freundlich equation to the correlation of data relating to the distribution of a solute between any two phases (gas-liquid, liquid-liquid, gas-solid, or liquid-solid) is suggested. This equation is readily adaptable to the usual methods of calculation involved in absorption and extraction.

541.123.2 2443

The molecular condition of phenols in benzene solution. DAVISON, J. A. J. Amer. Chem. Soc., 67, pp. 228-233, Feb., 1945.—From determination of the freezing points of solutions of Ph3CH in C6H6 in an apparatus designed for the thermo-electric measurement of the freezing points of dilute solutions (0.002-0.1 M) the cryoscopic constant of the solvent was calculated as  $5.088 - 0.0548 \Delta T$ . The freezing points of solutions of PhOH, o- and p-CH3C6H4OH, and m- and p-NO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OH in C<sub>6</sub>H<sub>6</sub> were found. PhOH was shown to form solid solutions in C<sub>6</sub>H<sub>6</sub>, and the partition coefficient of the solute between solid and liquid C6H6 was calculated to be 0.41. Solutions of PhNO<sub>2</sub> obeyed the ideal law to  $\pm 2\%$ below 0.1 M.; there was no evidence of solid solution formation. The deviations from ideal behaviour shown by o-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>OH, p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>OH, pNO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OH, and m-NO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>OH increased in the order given. Equilibrium constants for dimer formation by these substances, and by 2:4-(NO<sub>2</sub>)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>OH, which is chelated, were calculated.

W. R. A.

541.123.2: 536.7 see Abstr. 2283

541.123.3

2444

Liquid-vapour equilibrium in methane-ethyleneisobutane system. BENEDICT, M., SOLOMON, E., AND RUBIN, L. C. Industr. Engng Chem., 37, pp. 55-59, Jan., 1945.—Liquid-vapour equilibrium data are presented for 37.78° and 71.11°C, and 34.02 and 68.05 atm. The data agree with previous predictions. [Abstr. 871 (1943)].

541,123,33 : 532,733 see Abstr. 2191

541.123.5

2445

Some properties of organic ion-exchange materials. HOLMES, E. L. J. Soc. Dy. Col., Bradford, 61, pp. 39-40, Feb., 1945.—Two types of reactive synthetic resins are used as cation- and anion-exchangers-the former owing their property to the presence of sulphonic-groups and the latter to basic N groupings. Materials other than synthetic resins have not met with the same success. For utility, the main criterion is insolubility especially for use in water treatment where it is essential both for the purity of the treated water and for the life of the materials themselves. Materials which contain replaceable H atoms obtain a fresh supply of ions by treatment with dilute acid solution. The anion-exchanger can have renewal by treatment with dilute caustic, carbonate or bicarbonate Natural waters passing through the two exchangers have their electrolyte content reduced by at least 95%, and treated water has practically the quality of distilled water. н. н. но.

541.123.5 2446

Liberation of H+, Al+++ and Fe+++ ions from hydrogen clays by neutral salts. MUKHERJEE, J. N., AND CHATTERJEE, B. Nature, Lond., 155, pp. 268-269, March 3, 1945.

541.123.7 : 518.3 2447

Representation of equilibrium constant data. Scheibel, E. G., and Jenny, F. J. Industr. Engng Chem., 37, pp. 80-82, Jan., 1945.—A nomograph gives the equilibrium-constant data for hydrocarbon mixtures over a wide range of temp, and pressures. Its construction is empirical, with the temp. scale based on the liquid fugacity of a reference substance and the pressure scale based on its vapour fugacity.

Kinetics of wood saccharification. Hydrolysis of cellulose and decomposition of sugars in dilute acid at high temperature. SAEMAN, J. F. Industr. Engng Chem., 37, pp. 43-52, Jan., 1945.-A method is described for hydrolyzing small samples of cellulosic materials in glass bombs by heating with direct steam in a rotating autoclave. The decomposition rates of glucose and other sugars in dilute acid were investigated, and the decomposition found to follow the laws of a first-order reaction. The energy of activation of the decomposition of glucose appeared to be independent of the acid conc. over the range 0.4-1.6% H<sub>2</sub>SO<sub>4</sub>, and averaged 32 800 cal. The rate of reaction increased 125% with a 10% rise in temp.

2441

between 170° and 190°C. The energy of activation in the hydrolysis of Douglas fir was independent of the acid concentration and averaged 42 900 cal.

541.124 244

Quantitative determination of methylglyoxal, and the mechanism of its reaction with hypoiodite solution. Woo, S. C., AND CHANG, S. T. J. Chem. Soc., pp. 162-165, March, 1945.

541,124: 541.64: 541.145 see Abstr. 2491

541.124: 577.15

Energy relationships in enzyme reactions. FRUTON, J. S., BALL, E. G., BERGMANN, M., KALCKAR, H. M., MEYERHOF, O., AND SMYTHE, C. V. Ann. N.Y. Acad. Sci., 45, 9, pp. 357-435, 1944.—The series of papers is the result of a conference on Energy Relationships in Enzyme Reactions held by the Section of Physics and Chemistry of The New York Academy of Sciences, February 11 and 12, 1944. Introductory remarks to the Conference on Energy Relationships in Enzyme Reactions, Fruton, J. S.; Energy relationships of the oxidative enzymes, Ball, E. G.; Energy relationships in glycolysis and phosphorylation, Meyerhof, O.; The function of phosphate in enzymatic syntheses, Kalckar, H. M.; The significance of coupled reactions for the enzymatic hydrolysis and synthesis of proteins, Bergmann, M., and Fruton, J. S.; Some enzyme reactions of sulphur compounds and their possible interrelationship, Smythe, C. V.

541.124:676.1 245

Isolation and utilization of lignosulphonic acids. Pepper, J. M. Pulp. Pap. Mag. Can., 46, pp. 83-91, Feb., 1945.—A review, with extensive bibliography, of recent work on the chemical mechanism of the sulphite pulping process for wood, the classification of the lignosulphonic acids, their industrial recovery by evaporation or precipitation, and their properties and utilization. The last includes uses as a fuel, and as a source of vanillin, tanning agents, dyes, binding materials, lignin plastics, emulsifying agents, lubricants, decolorizing agents, bleaching agents, hydrogenation products, and fertilizers.

J. G.

541.124/.127 2452

Kinetics of the base-catalyzed reaction of pseudo-I-menthyl I-o-benzoylbenzoate with methanol. Schaefgen, J. R., Verhoek, F. H., and Newman, M. S. J. Amer. Chem. Soc., 67, pp. 253-257, Feb., 1945.—Polarimetric measurements of the reaction in the presence of methylate ion to form methyl o-benzoylbenzoate shows it to be pseudo-first order, and specifically catalyzed by methylate ions. The activation energy is 11 000 g.cal. No primary salt effect was detected. Fission of the pseudo ester takes place between the alcoholic oxygen and the carbonyl carbon of the acid. A mechanism for the reaction is postulated.

W. R. A.

541.124/.127: 541.144.7 see Abstr. 2488

541.124/.128: 535.243 see Abstr. 2223

541.124.7: 541.135.2 see Abstr. 2480

541.124.7: 541.145 see Abstr. 2492

541.124.7: 541.57: 541.144.7 see Abstr. 2489

541.126 2453

The spontaneous combustion of hay. FIRTH, J. B.,

AND STUCKEY, R. E. J. Soc. Chem. Ind., 64, pp. 13-17, Jan., 1945.

541.126 2454

Mechanism of the oxidation of coal. Jones, R. E., AND TOWNEND, D. T. A. Nature, Lond., 155, pp. 424-425, April 7, 1945.

541.126: 662.4 2455

Determining the initiating efficiency of detonators. GRANT, R. L., AND TIFFANY, J. E. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 13-19, Jan., 1945 .-The miniature-cartridge test is based on the principle that the ability of a detonator to initiate an insensitive explosive constitutes the best means of measuring its initiating efficiency. The experimental technique combines the features of the sand test and the TNTiron-oxide insensitive-powder test. A unique operation is the subtraction of a detonator blank. This achieves a measurement of actual resultant effect of detonator in producing a greater or lesser degree of detonation in an insensitive explosive. Results of tests are presented as curves for Hg(CNO)2-KClO3 (80-20) detonators, and tetryl-base detonators. From these, a procedure for a routine test is developed and outlined.

541.127 2456

The thermal decomposition of mercuric fulminate. NARAYANA, P. Y. Curr. Sci., 13, pp. 313-315, Dec., 1944.

541.127 2457

Kinetics in acid media: the correlation of reaction rates with acidity function. Deane, C. W. J. Amer. Chem. Soc., 67, pp. 329-331, Feb., 1945.—Acidity function data for fuming H<sub>2</sub>SO<sub>4</sub> are correlated with kinetic data for the decompositions of certain organic acids in this medium.

W. R. A.

541.127 2458

The mechanism of the hydrogen fluoride catalysed alkylation of toluene by t-butyl chloride. PEARLSON, W. H., AND SIMONS, J. H. J. Amer. Chem. Soc., 67, pp. 352-362, March, 1945.—The alkylation of PhMe by t-BuCl, catalyzed by HF, has been followed by the rate of increase of pressure in a special all-metal apparatus the inner surface of which is Au plated. With H<sub>2</sub>O, McOH, Et<sub>2</sub>O and hexamethylacetone as promoters, the rate of reaction increases with increasing conc. of promoter, but all give essentially the same effect at the same molar conc. The equations of Sprauer and Simons [Abstr. 2426 (1942)] are modified to correlate the rate measurements in any one experiment, but their equations and theory are essentially confirmed. Rate measurements and the effects of the promoters confirm the contribution of the amphoteric medium effect to the mechanism of the reaction. A one-step, condensed phase, catalyzed, and promoted reaction as the essential factor in the mechanism is shown to be satisfactory from kinetic reasonable 'from energy considerations. Mechanisms involving ions or free radicals as intermediates are shown to be untenable from energy considerations. W. R. A.

541.127 : 541.128 see Abstr. 2466

541.127: 541.128.1

Kinetics of the catalytic dehydrogenation of alcohols and glycols in the gaseous state. Neish, A. C. Canad.

J. Res. B, 23, pp. 49-69, March, 1945.—The catalytic dehydrogenation of several alcohols and glycols, in the gaseous state, was found to be a first-order process obeying the Arrhenius relation for variation of rate with temperature. Hence a large body of experimental data can be expressed by 2 constants that enable a ready comparison of different catalysts. A porous Cu catalyst was the most satisfactory. The fallacy of calculating the energy of activation from the temperature coefficient in this type of reaction is pointed out. Primary alcohols are dehydrogenated more slowly than secondary because the resulting aldehydes reversibly depress the activity of the catalyst, while methyl ketones have no measurable effect. Diacetyl acts as an aldehyde in this respect. The α-glycols form α-hydroxy aldehydes and ketones that depress the activity of the catalyst to a low but constant level. Rapid poisoning of the catalyst at constant temperature is a first-order process. A regular relation exists between the temperature of reduction and the activity of the catalyst.

541.127:541.57:536.66 2460

Rates of pyrolysis and bond energies of substituted organic iodides. II. BUTLER, E. T., MANDEL, E., AND POLANYI, M. Trans. Faraday Soc., 41, pp. 298-306, June, 1945.—A continuation of the account of the pyrolysis of organic iodides at low pressures and brief times of reaction [Abstr. 920 (1943)]. Under these conditions the rates are determined by the primary decomposition XI = X + I and thus characterize the bond energy C-I in various compounds XI. The paper contains data concerning the pyrolysis of cyclohexyliodide,  $\beta$ -phenylethyliodide, isopropyliodide, t-butyliodide, dichloroiodomethane, dibromoiodomethane,  $\beta$ -chloroethyliodide and iodoform. For the first two of these, the bond energy could be determined with considerable certainty, since the value of the activation energy was the same whether calculated as

 $Q = -RT^2 d \log_e k_1/dT$  or as  $Q^* = 2 \cdot 3RT(13 - \log k_1)$ where  $k_1$  is the mono-molecular rate constant.

541.127 : 542,929,4 2461

Alkaline permanganate oxidation of certain condensed cyclic compounds including coal. WARD, J. J., KIRNER, W. R., AND HOWARD, H. C. J. Amer. Chem. Soc., 67, pp. 246-253, Feb., 1945.—Apparatus has been devised and analytical methods have been worked out for, inter alia, following quantitatively the rate of reaction of condensed cyclic compounds with alkaline permanganate, permitting work to be done on samples of 500-800 mg. For comparison, a sample of a typical bituminous coal was investigated under identical conditions. The rate of oxidation diminished in the following order: tetrahydronaphthalene, bituminous coal, pyrene, naphthacene, diphenylene oxide, carbazole, decacyclene and triphenylene; the nearly saturated structures were least reactive of all. A qualitative correlation between the rates of oxidation and "double bond character" of certain polycyclic hydrocarbons was found. Significantly higher rates of oxidation were found than have been previously reported for some of these compounds, probably due to the excess permanganate present in the early stages of the reaction.

541.127.2:616.9

2462

The law of mass action in epidemiology. WILSON, E. B., AND WORCESTER, J. Proc. Nat. Acad. Sci., Wash., 31, pp. 24-34, Jan., 1945.—The law of mass action states that the rate at which an infection passes in a population is proportional to the product of the number of persons who are infectious and the number of persons who are susceptible to the infection. Various modifications of the law are considered and a mathematical analysis of these cases is given. This involves, for the most part, approx. solutions of various differential equations.

L. S. G.

541.128 2463

The metal-catalysed reaction between acetylene and hydrogen. II. Further experiments with nickel catalysts. Sheridan, J. J. Chem Soc., pp. 133-142, March, 1945.—[See Abstr. 350 (1945)]. The effect . of temperature on the reaction between C2H2 and H2 over a Ni-pumice catalyst was studied in the range 0-126°. Lowering the temperature increases the production of C<sub>2</sub>H<sub>4</sub> and (more markedly) C<sub>2</sub>H<sub>6</sub>; the yield of  $C_2$  hydrocarbons, in terms of the  $C_2H_2$  not recovered, is 31% at 126° and 65% at 0-3° for 1:1 gas mixtures. The ratio of the  $C_2H_2$  removed as other products to the H2 so removed increases as the temp. is raised. Temp. (0-91°) has little influence on the orders of the reactions occurring. With some not improbable assumptions, activation energies, uncorrected for desorption of gas, are deduced as follows: (i)  $C_2H_2 + H_2 + Ni \rightarrow C_2H_4 + Ni$ : 10.9 kg. cal. at  $0-126^\circ$ . (ii)  $C_2H_2 + xH_2 + Ni \rightarrow$  products other than  $C_2$  hydrocarbons + Ni: 14.5 kg. cal. at  $0^\circ$ , falling to 12.7 kg. cal. at  $126^\circ$ . The mechanisms of the reactions are discussed, and it is suggested that reaction (ii) is a surface chain reaction, initiated through a half-hydrogenated state in associatively chemisorbed acetylene.

541.128 2464

The metal-catalysed reaction between acetylene and hydrogen. III. Some effects of added substances on the reaction over nickel. Sheridan, J. J. Chem. Soc., pp. 301-305, May, 1945.—[See Abstr. 2463 (1945)]. Reaction over Ni on a number of carrier substances (chiefly oxides) differs little from that over Ni-pumice. No influence of CO<sub>2</sub> or N<sub>2</sub> is detected. More than ca. 3% of NO retards reaction and lowers the yield of C<sub>2</sub>H<sub>4</sub> slightly; liquids containing N<sub>2</sub> are formed. Some reduction of NO by H<sub>2</sub> probably occurs. O<sub>2</sub> retards reaction and raises the C<sub>2</sub>H<sub>2</sub> yield. It is removed partly as H<sub>2</sub>O, but also reacts with the catalyst, its effects persisting during several succeeding experiments. SiO<sub>2</sub>, TiO<sub>2</sub>, ThO<sub>2</sub>, ZnO, Cr<sub>2</sub>O<sub>3</sub> catalyse polymerization in C<sub>2</sub>H<sub>2</sub> alone at -200°.

541.128 2465

The metal-catalysed reaction between acetylene and hydrogen. IV. Reaction over platinum. SHERIDAN, J. J. Chem. Soc., pp. 305–311, May, 1945.— [See Abstr. 2464 (1945)].  $C_2H_4$  is not formed selectively from  $C_2H_2$  and  $H_2$  over Pt-pumice at 0–163°, though  $C_2H_2$  is more strongly adsorbed than  $C_2H_4$ , and at pressures > a value (depending on the pressures of hydrogen and ethylene present) inhibits  $C_2H_6$  production to a rate < that of  $C_2H_4$  formation. With 1:1 and 2:1  $H_2:C_2H_2$  mixtures some

6-30% of the C2H2 is not recovered, and a comparable amount of H2, are removed as products less volatile than C2 hydrocarbons. Previous kinetic data are confirmed and it is shown that lower temperatures favour C2H6 production in the early stages. The reactions (i)  $C_2H_2 + H_2 = C_2H_4(\to C_2H_6)$ (ii)  $C_2H_2 + H_2 \rightarrow \text{products other than } C_2 \text{ hydro-}$ carbons are probably both of approx. 1st order with respect to H<sub>2</sub>, and have similar heats of activation (that of (ii) being the higher by ca. 1-2 kcal.) in the region of 12 kcal. The differences between (i) and (ii) are qualitatively the same as for the corresponding reactions over Ni. The stronger adsorption of C2H2 than of C2H4 is confirmed by the fact that C2H4 added to the system acts primarily as a diluent.

541.128: 541.127 2466

The catalytic hydrogenation of the benzene nucleus. I. The hydrogenation of phenyl-substituted aliphatic acids. Smith, H. A., Alderman, D. M., and Nadig, F. W. II. The hydrogenation of benzene and monoalkylbenzenes. III. The hydrogenation of polymethylbenzenes. SMITH, H. A., AND PENNEKAMP, E. F. H. J. Amer. Chem. Soc., 67, pp. 272-281, Feb., 1945 .-I. The kinetics of the catalytic hydrogenation of phenyl-substituted acids were studied in AcOH solution, using Adam's Pt catalyst. The reaction rate is first order with respect to H pressure, zero order with respect to the concentration of phenyl-substituted the rate of catalytic hydrogenation are discussed, II. The relative rates of hydrogenation of mono-alkyl benzenes are compared with those for the series of phenyl-substituted acids, and also with those for a series of C<sub>6</sub>H<sub>6</sub> hydrocarbons previously examined. III. Increased Me substitution in polymethylbenzenes decreases the rate of hydrogenation of the nucleus; the position of the substitution also affects the rate, and for compounds with the same number of Me substituents, symmetrically substituted derivatives react most rapidly. W. R. A.

541.128 : 548.74 : 548.73 see Abstr. 2592

541.128 : 621.891.2 2467

Factors causing lubricating oil deterioration in engines. Laboratory evaluation. Burk, R. E., Hughes, E. C., Scovill, W. E., and Bartleson, J. D. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 302-311, May, 1945.—[Abstr. 2288 B (1945)].

541.128.1: 541.127 see Abstr. 2459

541.128.5

The poisoning of nickel hydrogenation catalysts by water vapour. Burford, W. B., and Frazer, J. C. W. J. Amer. Chem. Soc., 67, pp. 331-332, Feb., 1945.—Water vapour acts as a poison in the hydrogenation of  $C_2H_4$  with a Ni catalyst, the effect being noticeable down to  $3\times 10^{-4}$  mm. pressure of water vapour. The catalyst is reactivated by heating in  $H_2$  to 375°. W. R. A.

541.128.5 2469

The production of a poison for nickel catalysts in the absorption of oxygen by Fieser's solution. Burford, W. B., III, and Frazer, J. C. W. J. Amer. Chem. Soc., 67, pp. 341-342, Feb., 1945.—Fieser's solution No. 2, using an equivalent amount of KOH in place of the

NaOH specified, was used to remove oxygen from  $C_2H_4$  in the examination of the poisoning of Ni hydrogenation catalysts by water vapour. Highly erratic results were obtained due to the introduction of  $CS_2$  into the  $C_2H_4$ . The concentration of oxygen does not govern the amount of  $CS_2$  formed, and  $C_2H_4$  alone is without effect.

W. R. A. 541.132

The diffusion of ions in supporting electrolytes. DEAN, R. B. J. Amer. Chem. Soc., 66, pp. 31-35, Jan., 1945.—A supporting electrolyte is defined as the electrolyte (or mixture of electrolytes) initially present at uniform conc. throughout a system in which diffusion takes place. An equation is developed relating the integral diffusion coefficient of any ion to the conductivities of the supporting and diffusing electrolytes. The diffusion of chloride, Cu and citrate ions has been measured in various supporting electrolytes and the results agree with theory. The diffusion of ions is nearly independent of the relative ionic mobilities of the supporting electrolyte but depends almost entirely on the conductivity of the latter. When the supporting electrolyte has 10 times the conductivity of the diffusing electrolyte, an ion diffuses at a rate that still differs from its ideal rate, if uncharged, by 5% of the difference between the ideal rate and the rate in the absence of supporting electrolyte. For larger conductivity ratios the deviation will be proportionately less.

541.132

The temperature dependence of ionic dissociations. JENKINS, H. O. Trans. Faraday Soc., 41, pp. 138-140, March, 1945.—The experimental data for the electrolytic dissociation of many weak acids in water and dioxane-water mixtures were subjected to a theoretical analysis, using the equation

 $\log K = A/T + B \log T/T + C \log^2 T/T + D.$ 

This equation is satisfactory. Agreement with other equations was found for  $\Delta H^\circ$ ,  $\Delta G^\circ$  and  $\Delta S^\circ$ , but not for  $\Delta C_P$ . A similar type of curve is given when  $\Delta S^\circ$ ,  $\Delta H^\circ$  and  $\Delta C_P$  are plotted against the reciprocal of the permittivity of the medium.

541.132 2472

The extension of dissociation of salts in water. X. Dissociation minima. Davies, C. W. J. Chem. Soc., pp. 460-464, July, 1945.—In solutions in which the ionic strength increases sufficiently rapidly with increasing conc., ion-association will increase up to a critical conc., and thereafter will decrease. The degree of dissociation of the electrolyte will therefore have a minimum value. Evidence for this is found amongst aqueous solutions containing multivalent ions, and it is suggested that the effect will have an important bearing on the properties of colloid systems and non-aqueous solutions.

541.132 : 532.74 : 537.311 see Abstr. 2300

541.132 : 535.243 see Abstr. 2224

541.132 : 666.1

Energy additivity in glasses. Huggins, M. L., and Sun, K.-H. J. Soc. Glass Technol., 28, pp. 463-468, Dec., 1944.—Energies of dissociation into the component ions were computed for simple and complex oxides. From these, 51 energy constants are obtained, which may be used to compute approximate ionic

dissociation energies in crystals and glasses in which each metallic atom is surrounded by oxygens. The relation of these constants to the Periodic Table is shown and briefly discussed. Slight deviations from additivity in silicates and phosphates are related to the dependence of the energy on the number of oxygens in SiO<sub>4</sub> and PO<sub>4</sub> groups which are shared with other Si or P atoms.

541.132.3

pH of aqueous mixtures of potassium dihydrogen phosphate and disodium hydrogen phosphate at 0° to 60°C. BATES, R. G., AND ACREE, S. F. J. Res. Nat. Bur. Stand., Wash., 34, pp. 373-394, April, 1945.— A method is suggested for computing the pH of phosphate buffers from e.m.f. measurements of cells without liquid junction. Each of the 33 buffer solutions studied was prepared from equal molal quantities. The solutions were divided into 5 series with respect to the amount of NaCl added. The ratios of the molality of each buffer salt to that of NaCl in the 5 series were about 1, 2, 3, 8, and 10. The pH values were computed from measurements of cells with H electrodes and Ag-AgCl electrodes by a procedure that involves extrapolation of a function of the e.m.f. to zero concentration of NaCl. The values of the second dissociation constant of H3PO4 [Abstr. 1216 (1943)] were confirmed. The mean values of pK, the negative of the common log of the second dissociation constant, are given as a function of abs. temperature, T, by the equation

 $pK = 2.073 \cdot 0/T - 5.9884 + 0.020912T$ 

between  $T = 273 \cdot 16$  and  $T = 323 \cdot 16$  (0° to 50°C.). Equations are given to express the change of pH with molality of NaCl. The pH values from 0° to 60°C. of 8 phosphate buffers without chloride are listed. The densities of the buffers were determined. 541.132.3

Comparative liquid-junction potentials of some pH buffer standards and the calibration of pH meters. MANOV, G. G., DELOLLIS, N. J., AND ACREE, S. F. J. Res. Nat. Bur. Stand., Wash., 34, pp. 115-128, Feb., 1945.—By application of the equation pH  $=(E-E_{ref}-E_{j}/k)$  to solutions whose pH values were known accurately, the sum of the potentials of the reference electrode and of the liquidjunction potential  $E_{ref} + E_j$ , was obtained at 25°C. by the method of cells with liquid junction for 7 solutions suitable for standards of pH. Ag-AgCl electrodes immersed in saturated KCl solution were used. As  $E_{ref}$  remains constant when the buffer is changed, values for the differences in the liquidjunction potentials of various buffers in contact with saturated KCl solution were obtained from the data. These differences were then used to calibrate 7 Type 015 and 3 low-alkali-error glass electrodes of commercial manufacture. The average agreement between the true pH of the buffer-chloride solution (determined from cells without liquid junctions) and that read on various commercial pH meters when corrected for the difference in the liquid-junction potentials and the alkali error of the electrode was ±0.01 pH unit. The data also furnish a critical test of the consistency of the pH values assigned to the various buffer solutions recommended for the calibration of the pH scale and for checking pH meters.

541.132.3:620.193.92

2476

Effect of aeration on hydrogen-ion concentration of soils in relation to identification of corrosive soils. Romanoff, M. J. Res. Nat. Bur. Stand., Wash., 34, pp. 227-242, March, 1945.—62 air-dried soil samples taken from the N.B.S. soil-corrosion test sites were saturated and stored without air for almost 1 year. Significant changes in pH were observed in the majority of the samples and large changes in many of them. For the identification of areas corrosive to iron and steel, pH measurements of soils should be made on samples maintained in the natural field condition.

541.132.3:663.63 2477

Purification of water by use of synthetic ion-exchange resins: Using pH as a control. Kenworthy, A. L., AND HOWARD, J. N. Soil Sci., 57, pp. 293-294, April, 1944.—H2O purified by synthetic ion-exchange resins (e.g. Amberlite) compares favourably with H<sub>2</sub>O obtained from a copper still; the deionized H<sub>2</sub>O is comparable to redistd. H<sub>2</sub>O. Conductivity determinations clearly indicate that an ordinary pH meter may be used in the detection of the break-through point of the resins. The meter may also be used to regulate the regeneration and washing process after each cycle. Using conductivity determinations as a measure of purity the rate of the water flow through the resins had little effect on the purity of the deionized H<sub>2</sub>O. Germination of the sporangia of Phytophthora infestans indicated a continued adsorption of Cu after the cation exchanger is saturated. C. J. G.

541.134 2478

Hydrogen over-voltage in concentrated sulphuric and phosphoric acid solutions. De Béthune, A. J., And Kimball, G. E. J. Chem. Phys., 13, pp. 53-63, Feb., 1945.—The over-voltage for the discharge of H on a Hg cathode has been measured at 25°C. in H<sub>2</sub>SO<sub>4</sub> solutions in the range 0·125 to 9·65 M. In dilute solution, the over-voltage is independent of the acid conc., the cathode reaction being the discharge of a proton from water. At conc. > 2·83 M, the over-voltage decreases approximately as a linear function of the weight percentage. In this region the cathode reaction involves the H<sub>2</sub>SO<sub>4</sub> molecule. Preliminary results on H<sub>3</sub>PO<sub>4</sub> solutions also show a drop in the over-voltage as the concentration is increased beyond 1 M.

541.135 2479

Periodic phenomena at anodes. BARTLETT, J. H. Phys. Rev., 67, p. 268, April 1 and 15, 1945.—If a storage battery is connected through a resistance R to an electrolytic cell, an electric current may be made to flow through the cell. For a certain range of values of R and of the voltage across the cell, the current may oscillate continually and it may show a well-defined period.

541.135.2 : 541.124.7 2480

The kinetics of the electro-deposition of the azide ion. Stout, H. P. Trans. Faraday Soc., 41, pp. 64-75, Feb., 1945.—The anodic deposition of the azide ion from aq. solution at Pt, Pd, and Ir electrodes is an irreversible process giving a linear  $V/\log i$  curve with  $\alpha$  approx. = 1. The reaction is  $N_3 = 3/2 N_2 + e^-$ , and the reversible potential for unit activity of  $N_3$  is estimated to be  $-3\cdot3$  V against a normal H

electrode. Deposition occurs at measurable rates only at the high overpotential of 4 V. Increase in temperature causes a small but significant increase in  $\alpha$ , and the energy of activation is about 12 kcal/mole at a potential of 0.81 V against saturated calomel. The rate of reaction at a given potential varies with electrode metal, and with conc. of solution, but is independent of pH up to 9.4. At higher pH's,  $O_2$  is preferentially evolved.

541.135.5 2481

A microcalomel electrode for polarographic measurements. West, P. W., AND AMIS, E. S. Science, 101, pp. 71-72, Jan. 19, 1945.

541.135.5 2482

Membrane equilibria which involve only the ions of strong inorganic electrolytes. Sollner, K., and Gregor, H. P. J. Amer. Chem. Soc., 67, p. 346, Feb., 1945.—The use of recently developed collodion and protamine collodion membranes for the experimental study of Donnan membrane equilibria involving strong inorganic electrolytes is discussed. These membranes combine extreme ionic selectivity with great permeability for the non-restricted ion species. The ionic screening action of the membranes is primarily an electric and not a mechanical effect. Experimental results for K\*, Na\* and NH\* salts are given. W. R. A.

541.135.5 : 545.372 see Abstr. 2543 541.135.6 2483

The anodic behaviour of metals. I. Platinum. HICKLING, A. Trans. Faraday Soc., 41, pp. 333-339, June, 1945.—The initial build-up of anodic polarization at a smooth Pt anode over a wide range of conditions was investigated by an oscillographic method. The experimental results of previous workers are confirmed and extended, and 3 main stages in the polarization are distinguished corresponding to the ionization of H, the charging of a double layer, and the deposition of O at the electrode; it is suggested that the last process corresponds to the formation of a unimolecular layer of platinous oxide on the electrode surface. Attention is drawn to a singular anomaly which occurs in the presence of halides, when the anode potential rises directly to that for halogen evolution even where this is higher than the value at which oxygen can normally begin to deposit on the anode surface.

541.138 2484

Effect of substituents on the oxidation potential of ferrous ortho-phenanthroline complexes. Ewens, R. V. G. Nature, Lond., 155, p. 398, March 31, 1945. 541.14:545.8 2485

Photochemical analysis. Estimation of ferric salts. RAO, G. G., AND RAO, V. M. Curr. Sci., 13, pp. 317-318, Dec., 1944.

541.141:77.01 2486

Quantum theory of exposure tested extensively on photographic emulsions. SILBERSTEIN, L., AND TRIVELLI, A. P. H. J. Opt. Soc. Amer., 35, pp. 93-107, Feb., 1945.—Earlier [see Abstr. 298 (1943)] a formula was derived from the quantum theory of exposure for the characteristic curves of photographic emulsions. A detailed analysis has now been made of 6/700 curves for a great variety of emulsions under widely different development conditions. It is found

that the measured densities, D, for the total range of exposures applied, can be very closely represented either by a single or by two strictly quantic terms, that is to say functions of the exposure derived rigorously from the quantum theory under the assumption of a simple schematic form of size-frequency distribution in the emulsion.

A. H.

541.141.12 : 532.13

Effect of ultra-violet light on cellulose acetate and nitrate. Lawton, T. S., Jr., and Nason, H. K. Industr. Engng Chem., 36, pp. 1128-1130, Dec., 1944.— The effect on the viscosity of cellulose acetate and nitrate was studied in atmospheres of air, N<sub>2</sub>, and O<sub>2</sub>. In air and O<sub>2</sub> the degradation of these esters is accelerated. The effect of a N<sub>2</sub> atmosphere is to retard degradation. In all atmospheres the drop in viscosity for cellulose acetate was gradual; for cellulose nitrate the viscosity dropped more rapidly during the initial part of exposure and then levelled off. 541,144.7: 541,124/.127

Inadequacies in present knowledge of the relation between photosynthesis and the O<sup>18</sup> content of atmospheric oxygen. KAMEN, M. D., AND BARKER, H. A. *Proc. Nat. Acad. Sci., Wash.*, 31, pp. 8-15, Jan., 1945. 541.144.7: 541.124.7: 541.57

Bromination of hydrocarbons. I. Photochemical and thermal bromination of methane and methyl bromine. Carbon-hydrogen bond strength in methane. KISTIA-KOWSKY, G. B., AND VAN ARTSDALEN, E. R. J. Chem. Phys., 12, pp. 469-483, Dec., 1944.—The bromination of CH<sub>4</sub> was studied in the range 423-503°K., using a photo-electric photometer. A chain mechanism was found. Bromination of CH3Br is analogous but from 7.5 to 10 times more rapid in this temp. range. Thermal bromination, studied at 570°K., followed the same mechanism as photochemical reaction, except that Br atoms are produced thermally. The activation energy of photochemical bromination of CH<sub>4</sub> is 17.8 kcal/mole and that of CH<sub>3</sub>Br is 15.6 kcal/mole. Varying efficiencies of different molecules as third bodies in the homogeneous recombination of Br atoms are discussed. Configurations of activated complexes are assigned and shown to be reasonable by statistical mechanical calculations. Activation energies and other data give a value for the C-H bond strength in CH<sub>4</sub> of 102 kcal/mole at room temperature.

541.144.7 : 581.036

Effect of temperature on photosynthesis and respiration in red and loblolly pines. DECKER, J. P. Plant Physiol., 19, pp. 679-688, Oct., 1944.—Expts. were undertaken to discover whether the ratio, R, of photosynthesis to respiration in red pine is affected by high temp, and to account for the retarded growth of red pine in a forest in North Carolina. The rates of apparent photosynthesis and respiration were measured in seedlings of red and loblolly pine at 20, 30 and 40°C., light intensity of 4 500 ft. candles being used. The values for the two species were approx, the same at 30° as at 20°C, but decreased about 45% when the temp, was increased from 30° to 40°C. The value of R for both species was approx. 13 at 20°, 6.8 at 30°, and 3 at 40°C. Supplementary measurements of photosynthesis at a light intensity of 9 300 ft. candles and temps. of 19° and 32°C. showed no significant difference in the behaviour of the two species. R showed the same proportional decrease with increase in temp. in both species. There was no indication of a gross disturbance of the photosynthetic-respiratory balance of red pine at high temp, that might explain its retarded growth south of its normal range. The apparatus comprised two shoot chambers, a light source, three air sampling and analysis units and a flow-meter and valve for regulating the air-flow through the shoot chambers. The electrical components are a centrifugal fan, a heater, a thermo-regulated controller, and an electric blower which assured a continuous supply of air of nearly constant CO<sub>2</sub> content. 541.145: 541.124: 541.64

The mercury-photosensitized polymerization and hydrogenation of butadiene. Gunning, H. E., and Steacie, E. W. R. J. Chem. Phys., 12, pp. 484-493, Dec., 1944.—Collisions of butadiene mols. with Hg(3P1) atoms result in the formation of activated mols. of relatively long life, and these are mainly used up by the following reactions: (i)  $C_4H_6^* \rightarrow C_4H_4 + H_2$ ; (ii)  $C_4H_6^* + C_4H_6 \rightarrow 2C_4H_6$ ; (iii)  $C_4H_6^* + C_4H_6 \rightarrow (C_4H_6)_2$ . H atoms are formed when the H formed by (i) reaches a high enough conc. in the system to quench the Hg(3P<sub>2</sub>) atoms appreciably; polymerization then occurs by the reactions  $C_4H_6 + H \rightarrow C_4H_7$ , and  $C_4H_6 + C_4H_7 \rightarrow C_8H_{13}$  etc. N. M. B.

541.145 : 541.124.7 2492 The chain photolysis of acetaldehyde in intermittent light. HADEN, W. L., JR., AND RICE, O. K. J. Chem. Phys., 12, p. 521, Dec., 1944.—Erratum [see Abstr.

2922 (1942)]. 541.18:537,228.3

Electro-optical properties of colloids. II. SAKMANN, B. W. J. Opt. Soc. Amer., 35, pp. 66-85, Jan., 1945.—Investigations of the electric birefringence of colloidal solutions (monodisperse bentonite sols) in alternating fields show that, in general, the double refraction consists of a direct and an alternating part, both depending on field strength and frequency of the applied field, particle size, temp., and conc. of the solution. The direct birefringence, which can be positive or negative, can reverse its sign with a change of any of the 5 parameters mentioned. The measurements are interpreted by a relaxation theory which can satisfactorily explain all phenomena connected with the vibrating component, and the variations and the reversal of the direct birefringence with changes of the frequency, field strength, particle size, and temp. The changes and reversal of sign of the direct component with conc. are accounted for by assuming that the micelles are surrounded by a compressed water hull, the pressure of which depends on particle size and conc. Direct and alternating birefringence have different relaxation times [see Abstr. 2924 (1942)].

N. M. B. 541.18.041

Accuracy of mineral frequency analysis of sediments. ALLEN, P., AND WALDER, P. S. Nature, Lond., 155, pp. 173-174, Feb. 10, 1945. 541.18.043

A centrifuge tube agitator. REITEMEIER, R. F. Industr. Engng Chem. (Analyt. Edit.), 17, p. 267, April, 1945.

541.18.05

2496 A simple, rapid technique of preparing water-in-oil emulsions of penicillin, drugs and biologics. FREUND, J., AND THOMSON, K. J. Science, 101, pp. 468-469, May 4, 1945.

2497 . 541.182.021

Electron microscopic investigation of precipitates of cellulose nitrates. HAMBRAEUS, G., AND RANBY, B. Nature, Lond., 155, pp. 200-201, Feb. 17, 1945.

541.182.025 : 537.226.2 2498 Thixotropy and dielectric constant of printing inks. PARTS, A. Nature, Lond., 155, pp. 236-237, Feb. 24,

1945.

541,182.5 2499

Gel-forming derivative of wheat gluten. REITZ. H. C., FERREL, R. E., AND OLCOTT, H. S. Industr. Engng Chem., 36, pp. 1149-1151, Dec., 1944.-A product obtained from wheat gluten by the action of chlorosulphonic acid and pyridine or cold conc. H<sub>2</sub>SO<sub>4</sub> possesses, after neutralization, the property of absorbing rapidly 100-300 times its weight of cold water to form a firm, odourless, tasteless, and nontoxic gel.

541.182.5 : 542.952.6 see Abstr. 2534

541.182.6: 532.133 see Abstr. 2174 541,182.6 ; 537,362

2500

Comparison of the properties of freshly precipitated and heated aluminosilicates and alumina and silica gels and of clay minerals. RAYCHAUDHURI, S. P., AND DATTA, N. B. J. Phys. Chem., 49, pp. 21-31, Jan., 1945.—Nine kinds of aluminosilicate gels were prepared by mixing different proportions of SiO2 and  $Al(OH)_3$  sols  $(SiO_2 : Al_2O_3 \text{ ratios} = 2 : 1, 3 : 1, and$ 4:1), (1) by slowly adding Al(OH)3 sol to an excess of SiO<sub>2</sub> sol; (2) by mixing the two sols dropwise. The chemical composition (SiO<sub>2</sub>: Al<sub>2</sub>O<sub>3</sub> ratios), electro-osmotic charge, base-combining capacities, moisture contents at 50% r.h., and buffer curves were determined. These experiments were carried out also with electrodialysed alumina and silica gels and with the minerals montmorillonite, halloysite, kaolin, quartz, and bauxite. All these substances were subsequently heated in a steam autoclave for 12 hr. at 1 atm. pressure, and the physicochemical and electrical properties of the heated substances were determined. On heating, the precipitated aluminosilicates tend to acquire the properties of naturally occurring aluminosilicates. Mixing the colloidal solutions dropwise favours the formation of mineral structures.

541.182.6 : 539.26 see Abstr. 2392

541.182.6.046: 532.71

Note on the relationship between time of dialysis and the Burton-Bishop rule. TULLER, E. F., AND EBLIN, L. P. J. Phys. Chem., 41, p. 9, Jan., 1945.—Using NaCl and Na<sub>2</sub>SO<sub>4</sub> as coagulating electrolytes, it is found that those sols which have been dialysed until they are chloride-free conform to the Burton-Bishop rule [ibid., 24, p. 701, 1920]. Sols may conform to the rule before they have become chloride-free, and all sols exhibiting this unexpected combination of properties had been dialysed for 96 hr.

541.183.1 2502

The effect of the sub-phase upon the properties of physically adsorbed films. Jura, G., and Harkins, W. D. Proc. Amer. Phys. Soc., Chicago, Dec. 1 and 2, 1944. Abstr. in Phys. Rev., 67, p. 61, Jan. 1 and 15, 1945.—When the effective cross-sectional area of  $N_2$  on 100 solids is calculated, it is found that 3 max. are obtained at  $14\cdot 0$ ,  $15\cdot 2$ , and  $16\cdot 2$  Ų. In some cases there is a relation between the area exhibited and the composition or the crystal structure of the solid. The film of propyl alcohol on BaSO<sub>4</sub> is monomolecular, but polymolecular on anatase (TiO<sub>2</sub>) and quartz. The thickness of the film is related to the lattice constants of the solids. The transition from a gaseous to an intermediate film is first-order at 1 000 Ų per molecule for one solid while for another of the same crystal structure and particle size, the transition is second-order at 50 Ų per molecule.

541.183.1 2503

A theory of adsorption towers. BRIANT, R. C. Bull. Univ. Pittsburgh, 41, 4, pp. 38-47, 1945.—A mathematical treatment on the assumption that the fluid flowing through the tower has a definite unique value at each point and that this velocity is constant or changes slowly with time. Complications not directly of interest in tower operation are avoided; turbulence, flow irregularities, and the pressure drop in the tower due to flow are ignored. An equation is derived, suitable approximations are introduced, deductions of operational constants are shown, and applications to representative data are plotted and discussed.

N. M. B.

541.183.3 2504

Multiple adsorption from solutions. Walter, J. E. J. Chem. Phys., 13, pp. 229-234, June, 1945.—The general theory of the adsorption of solutes from solutions treated by a process similar to that used in chromatography is given. By assuming a chemical equilibrium between adsorbed material and the solution, the adsorption equations can be solved explicitly. A detailed discussion of the 2-component problem is given; the many-component problem and the extension of the theory to more general adsorption isotherms are considered.

541.183.526 2505

The sorption of hydrogen by poisoned copper. Temperature variation experiments. GRIFFIN, C. W. J. Amer. Chem. Soc., 67, pp. 62-64, Jan., 1945.— The sorption of H<sub>2</sub> with temperature variation on pure massive Cu was determined and compared with that of H<sub>2</sub> on the same Cu poisoned with a small quantity of CO. The effect of the poison is to prevent the solution of H<sub>2</sub> in Cu.

541.24 2506

Molecular weight of Palmer's β-lactoglobulin. Gutfreund, H. Nature, Lond., 155, p. 237, Feb. 24, 1945.—[See Abstr. 1868 (1944)].

541.24:535.435

An investigation of the determination of molecular weights of high polymers by light scattering. Doty, P. M., Zimm, B. H., and Mark, H. J. Chem. Phys., 13, pp. 159-166, May, 1945.—The determination of the molecular wts. of large molecules by measuring the turbidity of the solution and the change in the turbidity of the solution and the change in The apparatus, its calibration, and the technique used are described, with a comparison between molecular

wts. measured by this and other methods. The effect of polymolecularity is also presented.

541.265 2508

The effect of temperature on molecular parachors. GARDNER, T. S., SAMUEL, R. J. Chem. Phys., 13, p. 251, June, 1945.

541.265 : 535.324 2509

A study of organic parachors. VIII. A supplementary series of tertiary chlorides. SMART, K. O., AND QUAYLE, O. R. J. Amer. Chem. Soc., 67, pp. 21-23, Jan., 1945.—The refractive indices, densities, surface tensions and parachors for 9 supplementary tertiary chlorides are reported, at 3 temperatures. The predicted and experimental values of the parachors are in close agreement. A table is given for use in calculating the parachor of any tertiary chloride having normal alkyl groups.

541.42:632.94 2510

The stability of derris in insecticidal dusts. The solvent-powder effect. CAHN, R. S., PHIPERS, R. F., AND BRODATY, E. J. Soc. Chem. Ind., 64, pp. 33-40, Feb., 1945.—Oxidation of powders containing rotenone is greatly accelerated by addition of minor amounts of inert non-volatile liquid. Major factors controlling the rate of oxidation are the form and nature of the rotenone-containing material, the natures of the added liquid and of the solid phase, the amounts of rotenone-containing extract and added liquid and the ratios thereof, and the temperature. Oxidation is greatly decelerated by small amounts of strong acids, but is never totally suppressed; the amount of acid required for max. stability varies according to the nature of the preparation. The solvent-powder effect is held to be due to distribution of a solution of the extract over a large surface area. Stabilization by acid is believed to be due to the stability of derris, etc., increasing from alkalinity through neutrality to acidity; in neutral preparations oxidation, generally imperceptible in purely solid preparations or in bulk solutions, becomes important when accelerated by distribution of a solution over a large surface area. The phenomena are investigated by means of quantitative colour tests with particular reference to (i) the anti-louse powder, A.L. 63, Marks I and II, of the British Armed Forces, and (ii) the "conditioned dusts" of American agriculture. For these uses stabilization by acid has great practical value.

1 43 05 2511

Target chemistry of zirconium and yttrium; isolation of Zr89 in pure state. Kurbatov, M. H., and Kur-BATOV, J. D. J. Chem. Phys., 13, pp. 208-216, June, 1945.—Radioactive Zr89 obtained by bombardment of Y2O3 with deuterons, was separated without addition of stable Zr in quantities  $\simeq 10^{-10}$  gram. Oxychloride solutions of Zr89 mixed with inactive Zr were used for coagulation studies in the conc. range  $1.5 \times 10^{-4}$ - $1.5 \times 10^{-11}$  g.atom/l. It was found that the fraction coagulated increased with dilution under the const. conditions used (pH, 4.0; NaCl,  $2.5 \times 10^{-2}$  N; T, 25°C), to  $2.7 \times 10^{-6}$  g.atom/l. The log-log plot of Zr conc. (C) in solution against coagulated part (c) closely approaches a straight line with exponent n = 1.50 and k = 490. At  $1.5 \times 10^{-6}$ g.atom/l. and lower concs. the coagulated fraction reaches a max. and becomes const., k = C/c = 0.45.

2516

Within the range studied salt increases coagulation of Zr above  $10^{-5}$  g.atom/l.; but below  $10^{-6}$  g.atom/l. the salt effect its reversed. Increase in pH results in increase in the fraction coagulated (r) in all concs. studied. At a minute conc. r = (pH - K)/m, K being 1.85 with 0.05 N NaCl, and 1.58 with 0.005 N NaCl, and m being 3.15. The separation of  $Zr^{89}$  in minute quantities without addition of stable Zr is plausible if a Nernst distribution of Zr between coagulated part and solution is postulated.

541.454 : 541.63 see Abstr. 2520 541.454 : 541.636 see Abstr. 2522

541.5

Molecular constants and chemical theories. WHELAND, G. W. J. Chem. Phys., 13, pp. 239-249, June, 1945.—Topics from four papers by Samuel on the theory of valence are critically examined. Conclusions are that Samuel's bond-moments for molecules containing "pentacovalent N" are not as satisfactory as claimed, and provide no evidence in favour of the existence of pentacovalent N; that Samuel's eigenfunctions for structures with pentacovalent N are identical with the ones set up in the now conventional manner for structures without pentacovalent N but with semipolar double bonds instead; that the existence of antibonding orbitals seems to be theoretically necessary; and that several of Samuel's further conclusions are either based upon the use of words in unfamiliar senses, or else are erroneous.

541.5: 541.6

Structure of boron hydrides. Burawoy, A., Bell, R. P., and Longuet-Higgins, H. C. Nature, Lond., 155, pp. 328-329, March 17, 1945.

541.57: 536.66: 541.127 see Abstr. 2460

541.57:539.133

Bond moments of higher valence states. SAMUEL, R. J. Chem. Phys., 12, pp. 521-522, Dec., 1944.—In a previous paper additive bond moments were calculated for non-aromatic first-order molecules formed by a central atom in its maximal state of valency [see Abstr. 372 (1945)]. Whether the new values were really those of the semipolar double bonds remained doubtful; it is shown that this is not the case.

541.57: 539.133

III. Methyl cyanide Donor-acceptor bonding. addition compounds of boron trichloride and boron trifluoride. LAUBENGAYER, A. W., AND SEARS, D. S. J. Amer. Chem. Soc., 67, pp. 164-167, Feb., 1945 .-- V.d. measurements on CH3CN: BCl3 and CH3CN: BF3 show that dissociation of these compounds in the vapour phase is complete. Values of the heats of dissociation of CH<sub>3</sub>CN: BCl<sub>3</sub>(s) and CH<sub>3</sub>CN: BF<sub>3</sub>(s) are 33.4 and 26.5 kcal. respectively as calculated from these measurements; the calorimetrically determined heats of formation of CH3CN: BCl3(s) and CH<sub>3</sub>CN: BF<sub>3</sub>(s) from the gaseous components are 33.8 and 26.5 kcal. Cryoscopic measurements in C<sub>6</sub>H<sub>6</sub> show that in dilute solutions CH<sub>3</sub>CN: BF<sub>3</sub> is not appreciably dissociated. The dipole moment of CH<sub>3</sub>CN: BF<sub>3</sub> is approximately 5.8 debyes. W. R. A.

541,57: 539,133 see Abstr. 2366

541.57: 541.124.7: 541.144.7 see Abstr. 2489

6

On the structure of dimethyl ether-boron trifluoride. BAUER, S. H., FINLAY, G. R., AND LAUBENGAYER, A. W. J. Amer. Chem. Soc., 67, pp. 339-341, Feb., 1945.—Data previous obtained [Abstr. 2258 (1943)] are re-discussed. Assuming the dissociation of the  $(CH_3)_2O:BF_3$  complex at  $40^{\circ}C$ , previously considered negligible, to be about 50%, the structure in best agreement with the electron diffraction data has  $B-F=1:43\pm0.03$  Å,  $B-O=1.50\pm0.06$  Å,  $C-O=1.45\pm0.03$  Å. Electron diffraction examination of the  $CH_3CN-BCI_3$  complex showed it to be completely dissociated at 95°.

W. R. A. 541.6

The carbonato and bicarbonato pentammine cobalti ions. LAMB, A. B., AND MYSELS, K. J. J. Amer. Chem. Soc., 67, pp. 468-480, March, 1945.-The lack of structural significance of the water content of crystals of carbonato pentammine cobalti nitrate is shown. The dissociation constant of the bicarbonato pentammine cobalti ion is calculated from pH measurements on solutions containing mixtures of the bicarbonato and carbonato pentammine ions and their aquation products at 25°C, and of similar unaquated mixtures at 0°C. The values indicate a dipolar cationic structure for the carbonato pentammine cobalti ion and analogous ammines. The velocity constant for the aquation of the bicarbonato pentammine cobalti ion has been determined at 0°C and 25°C, and the equilibrium constant for the aquation of this ion calculated at 25°C. The carbonato pentammine cobalti ion appears to undergo a slow direct aquation. W. R. A.

541.6: 541.5 see Abstr. 2513

541.6:662.75:621.43.019

2518

The effect of the molecular structure of fuels on the power and efficiency of internal combustion engines. Kettering, C. F. Industr. Engng Chem., 36, pp. 1079–1085, Dec., 1944.—A lecture summary of the knock characteristics of fuels of various structures, expressed and plotted in terms of aniline equivalents. Long thin mols. knock badly and short compact ones of the same wt. only a little; the difference between them may amount to several-fold changes in the potential power output of an engine using them. The production and test of the efficient fuel "triptane" (2,2,3-trimethylbutane) is discussed in relation to engine design, and performance curves are given. N. M. B. 541.612

Distribution of chain lengths and compositions in copolymers. STOCKMAYER, W. H. J. Chem. Phys., 13, pp. 199-207, June, 1945.—The instantaneous distribution of chain compositions and chain lengths in vinyl copolymers is obtained in a simple form valid for long chains. The compositions of chains of a given length are normally distributed about the mean value, with a standard deviation which can be calculated from experimentally observable quantities. The distribution of chain lengths intimately resembles that for simple polymers.

541.63 : 541.454 252

Studies in stereochemistry. V. The effect of F-strain on the relative base strengths of ammonia and trimethylamine. VI. The effect of F-strain on the relative base strengths of ammonia and the methylamines. Brown, H. C. J. Amer. Chem. Soc., 67, pp. 374-380,

541.64:53

March, 1945.—V.—The relative base strength of NH<sub>2</sub> and NMe3 has been determined with Me3B, Et3B, (iso-Pr)3B and (tert-Bu)3B as the reference acids. The order NH<sub>3</sub> < NMe<sub>3</sub> observed with H acids and with Me<sub>3</sub>B is reversed with the other B acids as predicted from the effect of F-strain [Abstr. 1846 (1944)]. Improved methods for the preparation of Me3B and Et3B are described. The saturation pressures for the addition compounds of EtaB, (iso-Pr)3B and (tert-Bu)3B with NH3 and NMe3 are determined. VI.—The order of base strength NH<sub>3</sub> < Me<sub>3</sub>N < MeHN<sub>2</sub> < Me<sub>2</sub>NH observed with H acids and with Me3B is changed to Me3N< Me<sub>2</sub>NH < NH<sub>3</sub> < McNH<sub>3</sub> with (tert-Bu)<sub>3</sub>B as the reference acid, as predicted from the increased F-strain. The preparation of MeNH2. (tert-Bu)3B and Me2NH. (tert-Bu)3B are described. 541.634

Contributions to the stereochemistry of  $\gamma$ -carotene. Zechmeister, L., and Polgár, A. J. Amer. Chem. Soc., 67, pp. 108-112, Jan., 1945.—The cis-trans isomerization of  $\gamma$ -carotene,  $C_{44}H_{56}$  (from Minulus and Gazania flowers, lower melting form) was studied by several methods. Some stereoisomers are tentatively assigned configurations.

541.636: 541.454 2522 **B**-strain and base strength. Brown, H. C. J. Amer. Chem. Soc., 67, p. 503, March, 1945.—The C-P-C bond angle in Me<sub>3</sub>P is  $100 \pm 4^{\circ}$  and B-strain is therefore absent [Abstr. 1846 (1944)]. The order of base strength should be PH<sub>3</sub> < MePH<sub>2</sub> < Me<sub>2</sub>PH < Me<sub>3</sub>P; published data show the order to be PH<sub>3</sub> < MePH<sub>2</sub> < (Me<sub>2</sub>PH and Me<sub>3</sub>P). Saturation pressures at 25°C and 75°C for Me<sub>2</sub>PH. HCl and Me<sub>3</sub>P. HCl, as well as a competition experiment with a mixture of Me<sub>2</sub>PH, Me<sub>3</sub>P and HCl, give the order of strength Me<sub>2</sub>PH < Me<sub>3</sub>P. This is taken as strong support for the B-strain hypothesis. W. R. A.

Division of High-polymer Physics of the American Physical Society. J. Chem. Phys., 13, pp. 1-2, Jan., 1945.—List of papers presented at the inaugural meeting, on June 23-24, 1944.

541.64:539.42 Polymolecularity and mechanical properties of cellulose acetate. Sookne, A. M., and Harris, M. J. Res. Nat. Bur. Stand., Wash., 34, pp. 467-476, May, 1945.—The tensile strengths, ultimate elongations, and folding endurances of films prepared from a series of cellulose acetate fractions and blends were studied. When the mechanical properties are plotted against the intrinsic viscosities (or relative weightaverage degrees of polymerization), the results for the fractions and different blends fall on separate curves. In contrast, when the mechanical properties are plotted against the number-average degrees of polymerization, the results for the fractions and all of the blends fall approximately on a single curve for each property. The results are qualitatively consistent with the hypothesis that the mechanical properties of blends are the weight-averages of the properties of their components. The results emphasize the importance of the determinations of the number-average degree of polymerization in studying commercial polymolecular materials.

541.64: 541.124: 541.145 see Abstr. 2491 541.65: 537.533.73 see Abstr. 2318

The acidic constituents of Degras. A new method of structure elucidation. WEITKAMP, A. W. J. Amer. Chem. Soc., 67, pp. 447-454, March, 1945.—A new method of structure elucidation applicable to acids or amides with simple branched chains based on the number of transitions appearing in the solidification point curves of binary mixtures of the branched acid or amide with normal fatty acids or amides, is described.

541.68: 532.72: 532.13 see Abstr. 2168

A magnetic stirrer for continuous gas-flow apparatus. WINOKUR, M. Science, 101, pp. 49-50, Jan. 12, 1945.

542.231 : 542.74 2527

A combination gas-delivery tube and stirrer. RUSSELL, R. R., AND VANDERWERF, C. A. Industr. Engng Chem. (Analyt. Edit.), 17, p. 269, April, 1945.

542,231.7 2528 A chuck for glass tubing. Somerville, J. M.

J. Sci. Instrum., 22, pp. 114-115, June, 1945. 542.4: 533.52 see Abstr. 2195

542.48 2529

Determination of total nitrogen in proteins and their hydrolyzates. Improved method and apparatus. Jonnard, R. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 246-249, April, 1945.—A modified Kjeldahl distillation head is described.

542.48: 665.52: 545.727 see Abstr. 2545

542.66 2530 Modified construction of an automatic siphon. YARDLEY, V. A. J. Sci. Instrum., 22, p. 115, June,

542.67

Methods for the evaluation of analytical filter papers. SCRIBNER, B. W., AND WILSON, W. K. J. Res. Nat. Bur. Stand., Wash., 34, pp. 453-459, May, 1945.-Improved methods for rate of flow of water, retention of fine precipitates, and determination of ash, were developed. The use of complicated apparatus for measuring the rate of flow of water was avoided by applying Darcey's law to the ordinary conical filter. The use of pre-filtered water in making the measurement was found necessary for satisfactory reproducibility. For determining the ash content of paper, a tare crucible is used and the test specimen wetted and pressed into a compact wad to permit the use of as small a crucible as possible. The other tests were bursting strength of wet paper, thickness, weight per unit area, alpha cellulose, copper number, and acidity (pH).

542.74 : 542.231 see Abstr. 2527

542.74:621.369.2

Air drying by activated alumina. Engineering, 159, pp. 517-518, June 29, 1945.

542,929.4: 541.127 see Abstr. 2461

542.952.6 2533 Mechanism of vinyl polymerization. I. Rôle of

Mechanism of vinyl polymerization. 1. Role of oxygen. Barnes, C. E. J. Amer. Chem. Soc., 67,

2523

pp. 217-220, Feb., 1945.—An explanation is given for the frequently observed inhibiting action of oxygen on the photo-polymerization of vinyl compounds and some additional data are presented. It is shown that oxygen exerts a similar inhibiting action on thermal polymerization, and that both processes involve peroxide formation in preference to polymerization.

542.952.6: 541.182.5

Gel formation in addition polymerization. WAL-LING, C. J. Amer. Chem. Soc., 67, pp. 441-447, March, 1945.—An equation is deduced predicting the gel point in the addition polymerization of a mixture of mono- and bi-functional monomers. Gel points for the systems methyl methacrylateethylene dimethacrylate and vinyl acetate-divinyl adipate are determined under various conditions; the results agree reasonably with the equation in the presence of <0.2 mole% of bifunctional monomer. Results in the presence of more bifunctional monomer are explained qualitatively by considering the reaction mixture to consist of discrete swollen polymer molecules, the rate of diffusion of which is slow compared with the rate of polymer chain growth. On this basis, gelation is expected to occur when these swollen molecules have filled 13-23% of the reaction mixture. Experimental values of 25-46% agree with calculation.

W. R. A. 542.978 : 614.841 2535

The effect of chemicals in water solution on fire extinction. BRYAN, J., AND SMITH, D. N. Engineering, 159, pp. 457-460, June 8, and pp. 497-500, June 22, 1945.—Experiments are described on the extinction of standard fires under careful mechanically controlled conditions, with a method of recording continuously the loss in wt., by a jet playing on a rotating fire. Data for the relative eff. of solutions of a number of different chemicals are tabulated, plotted, and expressed as the traditional superiority factor, i.e. the ratio of water required to extinguish the fire to the amount of solution required to produce the same effect. NH<sub>4</sub> compounds (especially NH<sub>4</sub>H<sub>2</sub>PO<sub>4</sub> and (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>) show the most effective extinguishing properties. To assess the effectiveness in practice of different concs. of solution, and variation in effect at different stages of burning, a method of test employing a continuous application of liquid at varying rates is described. Data are discussed with reference to practical application. Results show that the true superiority is not the original "time superiority" factor, but is "volume superiority" factor, or ratio of vol. of water to vol. of solution necessary to extinguish the fire in the same time, and tends to a constant max, representing the max, potential fire fighting capacity of the solution, and this can be used for directly comparing relative efficiencies of different solutions irrespective of delivery. The potential firefighting capacity of water can be increased from 2 to 5 times by adding from 1.6-19% of NH<sub>4</sub>H<sub>2</sub>PO<sub>4</sub>.

N. M. B. 543,862,12 : 621,369.4 2536

Use of infra-red heat in determining aniline points. HOPKINS, H. T. Industr. Engng Chem. (Analyt. Edit.), 17, p. 269, April, 1945.

544: 669.14: 535.247.4 see Abstr. 2226

544.68 and seem not pulled a long about 2537

Spectrographic limit of detection of phosphorus, titanium, and zirconium in the direct current arc. NORMAN, D. P., AND JOHNSON, W. W. A. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 233-235, April, 1945.—A cored graphite electrode was used as the cathode of a conventional d.c. arc operated at 15 A and contained the weighed samples. The spectra were observed in the first order of a 3-m. grating spectrograph, dispersion 5.6 Å/mm., by a jumping-plate technique.  $\lambda 2553.28$  proved to be the most satisfactory line for the detection of P. The limit of detection of P, i.e. the smallest amount detectable with certainty, ranged from 0.05-0.80 microgram in 9 different standards containing from 0.0013-0.59% P. λ3371.45 was the most satisfactory for Ti; the limit of detection ranged from 0.04-4 micrograms in 12 samples containing from 0.0018-0.23% Ti. The optimum line for Zr was λ3391.98 of the ionized atom. The limit of detection of Zr varied from 0.5-4 micrograms in 6 samples containing from 0.0037-0.19% Zr.

545.31 2538

Sulphamic acid as an aid in the analytical electrodeposition of copper. SILVERMAN, L. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 270-271, April, 1945.

Application of the Ilkovič equation to quantitative polarography. BUCKLEY, F., AND TAYLOR, J. K. J. Res. Nat. Bur. Stand., Wash., 34, pp. 97-114, Feb., 1945.-Conditions of applicability of the Ilkovič equation to polarographic analysis were investigated for representative ions singly and in mixtures. Gelatin was used to suppress maxima found in current/voltage curves. Results obtained with a capillary of usual characteristics showed that a suppresser is necessary to secure agreement with the Ilkovič equation over any considerable drop-time range, and that the need increases with the drop-rate of the electrode and the dilution of reducible ion. A criterion of diffusion control satisfactory for classification of polarograms by visual examination was found. The application of the Ilkovič equation to quantitative analysis was shown to be inadvisable unless it is definitely established that the experimental conditions are well within the range for insuring diffusion control. A logarithmic relation between critical drop-time and suppressor concentration was found.

545.37

Polarographic characteristics of vanadium in its various oxidation states. LINGANE, J. J. Amer. Chem. Soc., 67, pp. 182–188, Feb., 1945.—From both acid and ammoniacal media V\*\*\*\* undergoes stepwise reduction at the dropping Hg-electrode to V\*\*\* and finally to V\*\*. In 0.05 N H<sub>2</sub>SO<sub>4</sub> the diffusion current of the first stage starts at zero applied e.m.f., and the half-wave potential of the second stage is -0.97 V against the standard calomel electrode. In 0.1 N H<sub>2</sub>SO<sub>4</sub> V\*\*\*\* is reduced irreversibly to V\*\* and the poorly-developed wave has a half-wave potential of -0.85 V against the standard calomel electrode. Reduction of V\*\*\*\* from M solutions of H ions is practically completely masked by the reduction of H\*. V\*\*\*\* from its N NaOH or KOH solutions

shows no indication of a reduction wave, but a well-developed anodic wave is obtained, corresponding to oxidation to V\*\*\*\*\*, with a half-wave potential of -0.42 V against the standard calomel electrode. The anodic diffusion current is proportional to the concentration of V\*\*\*\*, and the shape of the wave corresponds closely with a reversible one-electron oxidation. Both the reduction of V\*\*\*\*\* to V\*\*\*, and the reverse oxidation, are practically reversible at the dropping Hg-electrode from an acid medium. In NH<sub>2</sub>SO<sub>4</sub> both waves are well-developed, with an average half-wave potential of -0.52 V against the standard calomel electrode. Reduction of V below the V\*\* state was not encountered.

W. R. A. 545.37

Polarographic determination of vitamin C in fruits and vegetables. GILLAM, W. S. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 217-221, April, 1945.

545,371 2542

Potentiometric determinations of H-ion concentration in hydrocarbon oils. GEMANT, A. J. Chem. Phys., 13, pp. 146-154, April, 1945.—Oils were measured that have attained high electrical conductivity (a) through oxidation in the laboratory, (b) through deterioration in service, (c) through addition of certain compounds. The e.m.f. data are interpreted by comparison with those of lauryl sulphonic acid solutions in oil. The results indicate that the H ion concentration in the three types of oils is too low to account for the total observed conductivity. It is shown that the predominant mechanism of ion generation is different from the usual acidic dissociation. A bimolecular reaction leading to 2 organic ions is suggested and a quantitative relation following from the conductivity data established.

545,372 : 541,135.5

Determination of copper in copper proteins using the dropping mercury electrode. AMES, S. R., AND DAWSON, C. R. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 249-253, April, 1945.-A specific method for the determination of Cu in Cu proteins involves use of the dropping Hg-electrode after an acid extraction of the Cu. The base solution for analysis is an acid Na citrate buffer containing 0.005% fuchsin as a maximum suppressor. The half-wave potential at 25.0°C. for cupric ion in the base solution is - 0.18 V against the saturated calomel electrode. The diffusion coefficient at 25.0°C. of the cupric citrate complex is equal to  $0.43 \times 10^{-5}$ cm.2 sec.-1 Practical limits of the method are from 1 to 75-100  $\mu$ g./ml. of Cu in the base solution with an average deviation of  $\pm 3\%$ .

545.72: 545.8: 621.386.1: 621.791.75

A preliminary investigation of the constitution of mildsteel arc-weld deposits. SLOMAN, H. A., ROONEY, T. E., AND SCHOFIELD, T. H. J. Iron Steel Inst., 29 pp., July, 1945. Advance copy.—[Abstr. 2269 B (1945)].

545.727: 542.48: 665.52

Analysis of the light constituents in crude petroleum by low-temperature fractional distillation. ASKEVOLD, R. J., AND AGRUSS, M. S. Industr. Engng Chem. (Analyt. Edit.), 17, p. 241, April, 1945.

545.8: 541.14 see Abstr. 2485

545.8: 621.386.1: 621.791.75: 545.72 see Abstr. 2544 545.81 2546

Colorimetric estimation of aluminium in aluminum steel. Craft, C. H., and Makepeace, G. R. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 206–210, April, 1945.—A method is presented for estimating acid-soluble Alin steel by means of NH<sub>4</sub> aurintricarboxylate in the range 0·04–1·5% Al. The method is rapid, and the accuracy is equal to that of routine gravimetric procedures.

545.81 2547

Determination of iron in the presence of cobalt. Two-component colorimetric method. Brown, E. A. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 228-230, April, 1945.—A colorimetric thiocyanate method is given for the estimation of 0.07 to 0.5 mg. of Fe in the presence of variable amounts of Co up to 90 mg. A filter-type photometer with two colour filters is used to circumvent the interference of the Co ion colour. The method is rapid with an accuracy of  $\pm 3\%$ .

545.81 2548

Determination of cobalt in biological materials with nitrosocresol. Ellis, G. H., and Thompson, J. F. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 254-257, April, 1945.—A quantitative colorimetric procedure for the determination of from 0·02-25 µg. of Co in plant or animal tissues is described. The method depends on the formation of the coloured complex of Co with o-nitrosocresol which is extracted from the aqueous phase by ligroin and measured in a photoelectric colorimeter fitted with absorption cells 10 cm. in depth, requiring a volume of 1·8 ml.

545.81 2549
Colorimetric determination of minute amounts of nickel. PASSAMANECK, E. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 257-258, April, 1945.

545.81 2550

Acrolein determination by means of tryptophane. A colorimetric micromethod. CIRCLE, S. J., STONE, L., AND BORUFF, C. S. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 259-262, April, 1945.—Compared with previous methods, this method of analysis of acrolein gives a spectrally purer colour with a max. in the absorption curve, a colourless blank, and greater specificity.

545.81: 535.243 see Abstr. 2225

545.82 2551

The determination of nitrogen in ferro-alloys and other materials by direct Nesslerization without distillation. Newell, W. C. J. Iron Steel Inst., 5 pp., May, 1945. Advance copy.—The direct method, without distillation, for the colorimetric determination of N is further applied to the analysis of ferro-alloys by the addition of stabilizing colloids to the Nessler-NH<sub>3</sub> coloration.

545.82:669.14 2552

A microspectrographic method for the quantitative analysis of steel segregates. Convey, J., and Oldfield, J. H. J. Iron Steel Inst., 26 pp., May, 1945. Advance copy.

545.822 2553

Spectrochemical analysis of beryllium ores and commercial products. STROCK, L. W. J. Opt. Soc.

Amer., 34, p. 772, Dec., 1944.—Describes methods for rapid routine determination of Be in ores and commercial products. Ni is added as an internal intensity standard. The determination of impurities in BeO, Be-Cu alloy and Be metal and a new technique for establishing the major constituents of metallic alloy samples in a d.c. carbon are are described. J. W. T. W.

An application of spectrographic methods to chemical concentrations of trace elements in iron and steel analysis. Wolfe, R. A., AND FOWLER, R. G. J. Opt. Soc. Amer., 34, p. 772, Dec., 1944.—Describes a combined spectrographic and chemical method for the analysis of trace elements in steels, applicable to concs. below the sensitivity of either the ordinary chemical or spectrographic method.

J. W. T. W.

545.822

Spectrographic analysis of wrought copper alloys.
LEICHTLE, P. A., AND WICHROWSKI, A. F. J. Opt.
Soc. Amer., 34, pp. 772-773, Dec., 1944.—Discusses the spectrographic determination of trace elements and minor constituents in a wide variety of wrought Cu alloys. Tables of useful line pairs and approximate equal density values are given. [See Abstr. 173 (1945)].

J. W. T. W. 545.822

The spectrochemical analysis of copper base alloys. JAYCOX, E. K. J. Opt. Soc. Amer., 34, p. 773, Dec., 1944.—Describes the analysis of Cu base alloys for Al, Pb and Sn (0·30-15%), Fe, Mn and Ni (0·30-5%) and Zn (0·30-40%). A solution with a fixed conc. of metal is mixed with a fixed Cu(NO<sub>3</sub>)<sub>2</sub> solution and d.c. arc spectra are obtained for each sample and for a series of standard solutions. The amount of constituent element is found by measuring the ratio of the intensity of a line of the element to that of a Cu control line.

J. W. T. W.

545.824 2557 Chemical analysis by X-ray absorption. Lieb-

Chemical analysis by X-ray absorption. LIEB-HAFSKY, H. A., AND WINSLOW, E. H. Gen. Elect. Rev., 48, pp. 36-39, April, 1945.—A description of routine measurements.

545.83

Quantitative microchemistry in clinical laboratories. SOBEL, A. E. Industr. Engng Chem. (Analyt. Edit.), 17, pp. 242-245, April, 1945.

545.844 Separation and determination of aromatic and monoolefin hydrocarbons in mixtures with paraffins and naphthenes by adsorption. MAIR, B. J. J. Res. Nat. Bur. Stand., Wash., 34, pp. 435-452, May, 1945 .-The mixture to be analysed is introduced into the top of a column of SiO<sub>2</sub> gel and, when the liquid level just reaches the top of the gel, a suitable desorbing liquid, such as EtOH, is added. The desorbing liquid forces the hydrocarbon portion down the column, during which passage the hydrocarbon portion is fractionated according to the adsorbability of the various components. These components issue from the bottom of the column in the following order: Paraffin + naphthene, mono-olefin, and aromatic hydrocarbons. The analysis is made by determining the fraction of the total vol. constituted by each of these classes. Results of the analyses of 5 solutions containing aromatic and paraffin hydrocarbons, and 3 solutions containing aromatic, mono-olefin, and paraffin hydrocarbons are given. A procedure is outlined for determining the aromatic content of a straight-run petroleum distillate, as in the gasoline or kerosine fractions.

548.0:534.321.9 2560

Shear modes in piezo-electric crystal plates. BHAGA-VANTAM, S., AND SURYANARAYANA, D. Nature, Lond., 155, p. 171, Feb. 10, 1945.

548.0:535.32

Rodometric examination of quartz crystals. HOLTON, G. J. Electronics, 17, pp. 114-118 and 252, May, 1944.—The optical axis is approx. located by viewing through polarizing screens, and a surface normal to it (within 10°) is ground and etched in HF for 2-3 hr. A thin parallel beam of light projected through the crystal parallel to the optical axis, emerging from the ground "window," forms a characteristic star and spot pattern on a translucent screen. The position and sense of the X and Y axes, the "hand" of the crystal and the direction of the Z axis can be determined within about 1/4° from the pattern. Twinning is revealed. The apparatus is used to align the crystal axes in relation to a glass mounting plate, to which the crystal is cemented after adjustment. The mounted crystal can then be properly set for sawing. 548.0:535.43

Scattering of light in crystals. RAMAN, C. V. Nature, Lond., 155, pp. 396-397, March 31, 1945.— Discusses the fact that some theories of the solid state are at variance with experiment in that they assume, or claim to show, that the atomic vibration spectrum of a crystal in the infra-red region is continuous whilst experiment gives in some cases (e.g. calcite) a sharp line spectrum with no trace of a continuum. A. H.

548.0:536.76

Sub-crystalline changes of structure accompanying thermal transitions in Rochelle salt, and in potassium dihydrogen orthophosphate. UBBELOHDE, A. R., AND WOODWARD, I. Nature, Lond., 155, pp. 170-171, Feb. 10, 1945.

548.0:539.32 2564

Elastic constants of piezo-electric crystals. Zinc sulphide. BHAGAVANTAM, S., AND SURYANARAYANA, D. Proc. Indian Acad. Sci. A, 20, pp. 304-309, Nov., 1944.—An experimental method of detecting and measuring the resonance frequencies of piezo-electric crystal plates is described. The elastic constants of zinc blends have been obtained by this method as  $C_{11} = 10.79 \times 10^{11}$ ;  $C_{12} = 7.22 \times 10^{11}$  and  $C_{44} = 4.12 \times 10^{11}$  dynes/cm.<sup>2</sup> [see Abstr. 2565 (1945)].

548.0 : 539.32 : 534.321.9 2565

Elastic constants of crystals. A new method and its application to pyrites and galena. BHAGAVANTAM, S., AND BHIMASENACHAR, J. Proc. Indian Acad. Sci. A, 20, pp. 298-303, Nov., 1944.—A new method for determining the elastic constants of crystals, based on the transmission of longitudinal sound waves, is given, using a continuous ultrasonic spectrum and the Debye-Sears effect as a delicate test for detecting max. transmission. The method needs to be supplemented by at least one static-torsion experiment to obtain the

complete system of elastic constants. It has been applied to the cases of pyrites and galena. Voigt's results on pyrites are confirmed and values for the elastic constants of galena are given.

548.0: 539.56 see Abstr. 2433

548.1 2566

Crystal structure of barium titanate. Megaw, H. D. Nature, Lond., 155, pp. 484-485, April 21, 1945.

548.1:513.3 2567

Particle shape. WHITTAKER, E. J. W., SMITHSON, F., TOMKEIEFF, S. I. Nature, Lond., 155, pp. 331-332, March 17, and Corres., pp. 639-640, May 26, 1945.—A definition of criteria for distinguishing needle shapes from plates.

548.1:517.512.2 2568

Fourier transforms and structure factor. WRINCH, D. Phys. Rev., 67, p. 198, March 1 and 15, 1945.—This paper summarizes the properties of Fourier transforms with special reference to their use in crystal analysis. Every distribution has its own characteristic Fourier transform and it is shown how, once recorded, it can be used for the determination of its contribution to the structure factor of any crystal in which it is part of the unit pattern, since the Fourier transform of any lattice of points is also a lattice of points, namely the reciprocal lattice. Among those whose Fourier transforms are recorded are many simple point distributions, which yield the Fourier transforms of like atoms placed at such point sets when multiplied by the atomic scattering factor. Fourier transforms are also recorded for a number of mega-distributions including patterned cube and octahedral volumes and the corresponding shells and surfaces. The application of these transforms to the structure factors of small crystals is discussed. The Fourier transforms of a number of finite continuous distributions through cube and octahedral volumes and shells and on cube and octahedral surfaces are included.

and octahedral surfaces are included.

548.1:519.4

2569

Character tables for two space groups. Herring, C. J. Franklin Inst., 233, pp. 525-543, June, 1942.— The construction of character tables for the irreducible representations of space groups is fairly easy when each lattice point of the group possesses the full symmetry of the point group, but when this symmetry is lacking the construction is more laborious. Tables of the characters are given for the 2 most important groups not possessing the symmetry mentioned (i) the group  $D_{6h}^4$  (close-packed hexagonal) and (ii)  $D_h^7$  (diamond type). The general theory of representations is discussed.

548.1: 531.718 see Abstr. 2158

548.1:548.73

The crystal structures of metallic fluoberyllates, double fluoberyllates and sulphato-fluoberyllates. MUKHERJEE, P. L. Indian J. Phys., 18, pp. 148–158, June, 1944.—Crystals of the fluoberyllates of Rb, Tl, K and NH<sub>4</sub> were studied goniometrically and by X-rays. These crystals are isomorphous among themselves as well as with the crystals of the sulphates of the corresponding metals, both as regards crystal class and axial lengths. The space group of the crystals of the K compound was determined and

found identical with that of  $K_2SO_4$ . Crystals of the double fluoberyllates (a)  $Zn(NH_4)_2(BeF_4)_2 \cdot 6H_2O$ , (b)  $Co(NH_4)_2(BeF_4)_2 \cdot 6H_2O$ , and (c)  $Ni(NH_4)_2(BeF_4)_2 \cdot 6H_2O$  and also of the substance  $NiK_2BeF_4SO_4 \cdot 6H_2O$  are isomorphous among themselves as well as with the sulphates. The axial lengths of  $Ni(NH_4)_2(BeF_4) \cdot 6H_2O$  were measured and found identical with those of corresponding double sulphate.

548.1:549.211 2571

Are there four possible diamond structures? Lons-DALE, K. Nature, Lond., 155, p. 144, Feb. 3, 1945.— [See Abstr. 2637 (1944)].

548.2

A method of growing oriented sections of certain optical crystals. West, C. D. J. Opt. Soc. Amer., 35, pp. 26-31, Jan., 1945.—A method is described for growing extended crystal sections normal to a 3-fold axis from melts of NaNO<sub>3</sub>, and of alkali halides (KBr, KI). Such sections result when crystallization in the melt is started at a mica cleavage surface and is allowed to proceed in a direction normal thereto, the mica acting as an insoluble, infusible seed for the new crystal growth. The mica cleavage surface adsorbs from the melt a hexagonal net of alkali ions whose side is 5·17 Å, and this plane net constitutes the first layer of the new crystal growth.

548.24:553.621 2573 Terminology of interpenetrating twins in  $\alpha$ -quartz.

THOMAS, L. A. Nature, Lond., 155, p. 424, April 7, 1945.

548.32 2574

Isomorphic relationship between rubidium and thallium in igneous minerals. AHRENS, L. H. Nature, Lond., 155, p. 610, May 19, 1945.

548.526: 669.14: 539,378,3 see Abstr. 2405

548.7:536.77:531.19

A generalization of the quasi-chemical method in the statistical theory of superlattices. YANG, C. N. J. Chem. Phys., 13, pp. 66-76, Feb., 1945.—The quasichemical method in the investigation of the equilibrium distribution of atoms in the pairs of neighbouring sites in a superlattice is generalized by considering groups containing large numbers of sites. The generalized method may be used to obtain successive approximations of the free energy of the crystal. Integration is avoided by introducing a Legendre transformation. To analyse the fundamental assumption underlying the method more closely, the number of arrangements of the atoms for given long-distance order is calculated, and the hypothesis of the non-interference of local configurations is discussed. The method is applied to the calculation of the free energy in the different approximations discussed, including Bethe's second approximation and a simple approximation for the face-centred cubic crystal Cu<sub>3</sub>Au.

548.73 2576

On the origin of extra spots in Laue photographs. SIRKAR, S. C., AND BISHUI, B. M. Proc. Nat. Inst. Sci. India, 8, 2, pp. 217-231, 1942.—The experimental results published by previous workers and those obtained by the authors on the properties of the extra spots in Laue photographs are discussed in the light of the theories of diffuse scattering and of quantum reflection. The length of the streamers and the

structure of the extra spots observed in the case of diamond with particular glancing angles of incidence cannot be explained by the theory of quantum reflection (Raman and Nilakantan), but the details can be explained by the theory of diffuse scattering (Zachariasen). In the case of diamond, the observed directions of maxima also agree with Zachariasen's theory and the observed influence of temperature is given satisfactorily by the wave-mechanical theory (Born and Sarginson). The ratio of the intensity of diffuse scattering to that of Laue spot due to the TT plane of diamond increases with the increase of thickness of the crystal. These results are contradictory to the theory of quantum reflection and also to the results obtained by Venkateswaran in the case of NaCl.

548.73

Photography of crystal structures. Huggins, M. L. J. Chem. Phys., 12, p. 520, Dec., 1944.—Bragg's method of photographic summation of Fourier series by addition of the proper patterns of light and dark bands can be improved in accuracy and speed by the use of a previously prepared set of masks [see Abstr. 460 (1941)]. An improved set of 316 masks is reported, with which electron density photographs of crystal structures and Patterson summations are easily and rapidly made from the appropriate X-ray data.

548.73

A "reference" precipitated tricalcium phosphate hydrate. MACINTIRE, W. H., PALMER, G., AND MARSHALL, H. L. Industr. Engng Chem., 37, pp. 164-169, Feb., 1945.—Tricalcium phosphate hydrate, for use as a reference substance, is prepared by the slow addition of concentrated H<sub>3</sub>PO<sub>4</sub> to lime-concentrated sucrose solution. Stability in composition and reproducibility of product are established by X-ray and chemical analysis. Differentiation between this substance and hydroxyapatite may be effected by comparison of their 900°C. calcines.

J. T.

548.73 2579

The influence of absorption on the shapes and positions of lines in Debye-Scherrer powder photographs. TAYLOR, A., AND SINCLAIR, H. Proc. Phys. Soc., Lond., 57, pp. 108-125, March, 1945.—Simple graphical methods of determining the basic line-contours for powder diagrams taken in cylindrical Debye-Scherrer cameras are described. From these contours the absorption factor may be obtained for any given Bragg angle and dilution of the specimen. The contours also enable the position of the line peaks to be calculated for different sets of experimental conditions. This, in turn, opens up the possibilities of new types of extrapolation curves for the accurate determination of lattice parameters.

548,73

On the determination of lattice parameters by the Debye-Scherrer method. TAYLOR, A., AND SINCLAIR, H. Proc. Phys. Soc., Lond., 57, pp. 126-135, March, 1945.—The types of systematic error arising in the determination of lattice parameters by the use of Debye-Scherrer powder diagrams are discussed. The various extrapolation methods are reviewed, and it is shown how a consideration of the absorption factor and geometry of the focal spot lead to the most

satisfactory forms of extrapolation curve. It is shown how the absence of specimen eccentricity enables perfectly linear extrapolation curves to be drawn, thus allowing fullest use to be made of low-angle reflections, whereby the highest accuracy in parameter determination may be achieved.

548.73

The X-ray investigation of copper dl- $\alpha$ -aminobutyrate. Stosick, A. J. J. Amer. Chem. Soc., 67, pp. 362-365, March, 1945.—The unit cell is monoclinic with  $a=11\cdot09$  Å,  $b=5\cdot06$  Å,  $c=9\cdot45$  Å, and  $\beta=87^{\circ}$  51'. The probable space group is  $C_{2h}^5-P_{c}^{2_1}$  and there are two molecules per unit cell. The existence is shown of coplanar co-ordination of carboxyl O atoms and amino N atoms around the Cu atoms. Patterson and Fourier projections are employed.

J. T. 548.73

The crystal structure of nickel glycine dihydrate. Stosick, A. J. J. Amer. Chem. Soc., 67, pp. 365-370, March, 1945.—The unit cell is monoclinic with  $a=7\cdot60$  Å,  $b=6\cdot60$  Ă,  $c=9\cdot63$  Å and  $\beta=63^\circ25'$ . The probable space group is  $C_{2h}^5-P_c^{2_1}$  and there are two molecules per unit cell. The structure consists of slightly distorted, octahedrally co-ordinated complexes of two glycine residues and two water molecules about the Ni atoms. Patterson and Fourier projections are employed.

548.73 2583 X-ray crystallography of kojic acid. Fox, A. H.

Nature, Lond., 155, p. 397, March 31, 1945.

The validity of the pole figure. Decker, B. F. J. Appl. Phys., 16, pp. 309-310, May, 1945.—A [110] pole figure for cold-rolled and annealed silicon steel, constructed from data obtained with an integrating X-ray camera, is compared with a stereographic projection of [110] poles for 100 individual grains in the same samples. From this comparison, it is concluded that pole figures are accurate enough to determine preferred orientations when the exact percentage of crystals in each orientation is not

Some techniques for the growth and preparation of micro-specimens for X-ray diffraction studies. Fankuchen, I. Phys. Rev., 67, p. 197, March 1 and 15, 1945.—This paper presents techniques for the recrystallization on a micro-scale of materials for X-ray study from both solution and the melt. Techniques are also presented for mounting such specimens both in air and sealed in capillary tubes. The use of a micro-camera for the study of local variation in structure was discussed.

548.73 2586

An experimental investigation of extrapolation methods in the derivation of accurate unit-cell dimensions of crystals. Netson, J. B., AND RILEY, D. P. Proc. Phys. Soc., Lond., 57, pp. 160-177, May, 1945.— Measurements on X-ray photographs of cylindrical specimens of different absorption and thickness taken in a camera without eccentricity show that the absorption error in the apparent unit-cell dimension  $a \propto \cos^2\theta(\csc\theta+\theta^{-1})$ . The plot of a against

 $\frac{1}{2}\cos^2\theta(\csc\theta+\theta^{-1})$  is linear down to  $\theta=30^\circ$  for all 4 specimens used. The extrapolated values for a are in good agreement, and this extrapolation function is recommended in the case of data from well constructed cameras. Other extrapolation functions are also considered, and the effect of various sources of error discussed.

548.73:518.5

A method for calculating reciprocal spacings for X-ray reflections from a monoclinic crystal. BOOTH, A. D. J. Sci. Instrum., 22, p. 74, April, 1945.— The device consists of 2 perpendicular scales graduated in mm. upon one of which the extremity of a similarly divided hypotenuse scale can be set at any distance from the origin.

548.73:531.743

The two-circle goniometer in the X-ray laboratory. TUNELL, G. Phys. Rev., 67, pp. 197-198, March 1 and 15, 1945.—The principal uses of the two-circle goniometer in the X-ray laboratory are the orientation and adjustment of faceted crystals for X-ray diffraction photographs and the preliminary determination of axial elements. There are 3 types of two-circle goniometer; the Goldschmidt, the Czapski, and the Fedorov. None of these is available in the U.S. at present. If one is to be manufactured, it is imperative that the adjustment head be made interchangeable with the Weissenberg, oscillation, and Laue cameras on the market.

548.73: 535.312: 549.211: 537.531 see Abstr. 2308-2310

548.73:537.531:535.34 2589

Breadths of X-ray diffraction lines and mechanical properties of some cold-worked metals. MEGAW, H. D., AND STOKES, A. R. J. Inst. Met., 12, pp. 279–289, June, 1945.—The breadths of X-ray diffraction lines from cold-worked Fe, Ni, Cu, Ag, Al and Pb were measured, and shown to agree with those calculated from the mechanical properties of the metals on the assumption that the crystals are stressed up to the ultimate tensile limit.

548.73: 539.215.4 2590

X-ray diffraction examination of  $\gamma$ -alumina. Jellinek, M. H., and Fankuchen, I. Industr. Engng Chem., 37, pp. 158-163, Feb., 1945.—The results of the controlled heating of  $\gamma$ -alumina are investigated by low- and high-angle scattering of X-rays. Heating is found to increase both crystallite and particle size. The size of the cubic unit cell is given as 8.4 Å. J. T. 548.73: 539.26 see Abstr. 2393

548.73:539.388.1 2591

An X-ray diffraction study of the fatigue in metals at high stresses. BARKOW, A. G. J. Appl. Phys., 16, pp. 111-120, Feb., 1945.—Results are given of an X-ray diffraction study of 14 specimens of an Al alloy, 14 ST, at various cycles and stresses. A description is given of the preparation of the specimens and of the back reflection camera used [Abstr. 2393 (1943)]. The specimens were fatigued at stresses ranging from 25 000 lb./in². to 40 000 lb./in². L. s. G.

548.73 : 548.1 see Abstr. 2570 548.73 : 548.74 : 541.128

A review of the application of X-ray and electron diffraction methods to contact catalysis. MILLIGAN,

W. O. Phys. Rev., 67, p. 197, March 1 and 15, 1945 .-The results of X-ray and electron diffraction examination of contact catalysts is reviewed from the point of view of (a) identification of compounds in catalysts: (b) compounds formed during treatment or use; (c) small angle scattering; and (d) relation between catalytic activity and (1) lattice constants, (2) intensities, (3) crystal size, (4) orientation. mechanism of the formation of active catalyst is discussed from the point of view of (a) protective action, (b) solid solutions, (c) amorphous and crystalline states, (d) order and disorder, (e) distortion of crystal lattices. The state of absorbed substances and the potentialities of electron microscopic examination of catalysts are considered briefly. A bibliography of 135 references is included.

548.73:576.858.8

2593

X-ray crystallographic measurements on a single crystal of a tobacco necrosis virus derivative. Crow-FOOT, D., AND SCHMIDT, G. M. J. Nature, Lond., 155, pp. 504-505, April 28, 1945.—The crystals were 1-5 mm. across, and appeared to be of two forms: thick triclinic prisms and thin hexagonal or pseudohexagonal plates. X-ray data for the thickest crystal. which was of the first kind, showed that it was triclinic, with  $a = 179, b = 219, c = 243 \text{ Å}, \alpha = 87\frac{1}{2}^{\circ} \beta = 97\frac{1}{2}^{\circ}$  $\gamma = 971^{\circ}$  (reduced cell), space group P1. Photographs with the crystal "still" show numbers of X-ray reflections, and these, though individually unresolved. are characteristically arranged on the plate in series of concentric circles or ellipses. Oscillation photographs give further details of crystal structure. There are probably 4 units in the face-centred unit cell. The crystal structure suggests that the molecules are approximately spheres, and on this view the estimated mol. dimensions are = 80-100 Å radius in one direction. The sedimentation const. and other evidence indicate a mol. wt. of <1 850 000. N. M. B.

548.73: 621.386: 539.26 see Abstr. 2394, 2395

548.73:669.721.5

The crystal grain orientation in a rolled magnesium alloy. HARGREAVES, A. J. Inst. Met., 71, pp. 73-86, Feb., 1945.—The preferred orientation of the crystal grains in Elektron AM503 alloy sheet was examined by X-ray methods. In the surface of the sheet, the basal plane normals [0001] of the crystal grains are normal to the plane of the sheet, but in the centre of the sheet they tend to align themselves along one of two most probable directions inclined at approx. 15° on either side of the normal to the sheet and lying in the plane containing the rolling direction and the sheet normal. There is a gradual transition from one type of basal plane orientation to the other on passing from the surface to the centre of the sheet. The hexagon edges [1120] of the crystal grains are oriented parallel to the rolling direction in the surface of the sheet, but are oriented at random about the [0001] axes in the central regions of the sheet.

548.73:678.7

The structure of polyisoprenes. I. The crystal structure of geranylamine hydrochloride. II. The structure of  $\beta$ -guttapercha. Jeffrey, G. A. *Proc. Roy. Soc. A*, 183, pp. 388-404, *June* 18, 1945. *Trans. Faraday Soc.*, 40, pp. 517-520, *Nov.*, 1944.—I. The crystal structure and molecular dimensions are

established by an X-ray analysis employing threedimensional Fourier syntheses. The molecules have a trans configuration (similar to guttapercha) and lie parallel and end to end in pairs within an ionic framework where each N atom is equidistant from four Cl neighbours. The 2 isoprene units are planar and have normal interatomic distances, but are linked by a C-C bond markedly shorter than a normal single bond. This unusual feature is accompanied by a coplanar arrangement with adjacent C bonds. Tables are given of the calculated and observed structure factors. II. Present X-ray diffraction data are insufficient to distinguish fine details of molecular structure. A basis for a model of the  $\beta$ -guttapercha molecule is provided by the structure analysis of Part I, and this model differs considerably from that proposed by Bunn [Abstr. 1684 (1942)]. No evidence exists for the distortion of the methyl group out of the plane of the double bond previously ascribed to the molecule. For Part III see Abstr. 1420 (1945).

L. S. G. 548.73 2596

Photography of crystal structures by optical Fourier synthesis. Huggins, M. L. Phys. Rev., 67, p. 197, March 1 and 15, 1945.—[See Abstr. 460 (1941)]. A suitable set of band patterns was prepared on a roll of 35 mm. film. With this, typical electron-density and Patterson projections were made in less than 1 hr. The electron-density projections, so obtained, are true pictures of the unit of structure, magnified 108 or more times. The atoms are readily distinguished, as in the corresponding contour maps drawn from the results of the direct calculations.

548.74: 541.128: 548.73 see Abstr. 2592

549.211 : 535.312 : 548.73 : 537.531 see Abstr. 2308-2310

549.211: 548.1 see Abstr. 2571

549.514.51 : 621.92 2597

Control of silica dust hazard in quartz crystal cutting and grinding. Goss, A. E. J. Industr. Hyg. Toxicol., 26, pp. 208-209, June, 1944.—In operations using a diamond-edged circular saw and a recirculated soluble oil-water coolant liquid, in order to prevent the dissemination of silica dust into the atmosphere, the coolant liquid should be changed sufficiently frequently to keep the number of silica particles below  $3 \times 10^9$  particles/cm.<sup>3</sup> Manufacturing operations and sampling procedures are described. C. J. G.

55(764) 2598 Geology, climate and soils of Texas. BAKER, C. L. Proc. Trans. Tex. Acad. Sci., 27, pp. 181-187, 1943.

550:061.055.5

Geophysical Laboratory. ADAMS, L. H. Carnegie Instn Yearb., 43, pp. 21-22, 1943-1944.—Annual report.

550.3: 061.055.1 2600

Summary of the year's work, to June 30, 1944, Department of Terrestrial Magnetism, Carnegie Institution of Washington. FLEMING, J. A. Terr. Magn. Atmos. Elect., 49, pp. 245-250, Dec., 1944.

550.341 : 534.838.2 260

Earthquake-resisting structures. VALENZUELA, J. I. Engineer, Lond., 178, pp. 64-67, Jan. 26, 1945.—

Reference is made to disastrous earthquakes in Chile, Japan and San Francisco, and the consequent urge to improve the resistant qualities of buildings. Research on models and earthquake records has resulted in a new theory which takes into account the earth oscillations and also the mass and elasticity of buildings, thereby enabling the forces acting on the various members of a structure to be calculated. Earth tremors are propagated by longitudinal waves radiated from the hypocentre, followed by slower transverse waves; these are followed by the principal waves from the epicentre. The irregular motion of the epicentre causes great danger to buildings. Earth tremors may have periods from a few 1/100 sec. up to 10 sec., but the most destructive waves have a period of 1-2 sec. In relation to the foregoing the natural periods of different structures are examined, and a description is given of an elasticity meter designed and constructed in the University of Chile.

On the determination of magnetic vertical intensity, Z, by means of surface integrals. TAYLOR, J. H. Terr. Magn. Atmos. Elect., 49, pp. 223-237, Dec., 1944.—The main result of the present paper is a separation of Z, the normal component of a magnetic field, into parts  $Z_e$  and  $Z_l$  due to external and internal sources, respectively. The resolution is given in terms of surface integrals whose integrands involve the observed surface values X and Z of the field. The paper may be regarded as a sequel to an article by Vestine [Abstr. 1229 (1941)].

550.381 2603

List of geomagnetic observatories and thesaurus of values. VII. FLEMING, J. A., AND SCOTT, W. E. Terr. Magn. Atmos. Elect., 49, pp. 267-269, Dec., 1944.

550.381:061.055.5 2604
Department of Terrestrial Magnetism. FLEMING,

J. A. Carnegie Instn Yearb., 43, pp. 23-52, 1943-1944.—Annual report.

550.384 2605
Harmonic analysis of the annual variation of magnetic declination and horizontal intensity at Oslo.

netic declination and horizontal intensity at Oslo, 1843-1930. WASSERFALL, K. F. Terr. Magn. Atmos. Elect., 50, pp. 37-45, March, 1945.

550.384: 523.746

Solar and magnetic data, July to September, 1944,
Mount Wilson Observatory. Nicholson, S. B., and
Mulders, E. S. Terr. Magn. Atmos. Elect., 49,
pp. 277-278, Dec., 1944.

550.384: 523.746

Solar and magnetic data, October to December, 1944,
Mount Wilson Observatory. NICHOLSON, S. B., AND
MULDERS, E. S. Terr. Magn. Atmos. Elect., 50,
pp. 73-74, March, 1945.—[See Abstr. 2606 (1945)].

550.385 2608
Principal magnetic storms, July to September, 1944.
Terr. Magn. Atmos. Elect., 49, pp. 279-280, Dec.,

550.385 2609

Principal magnetic storms, October to December, 1944. Terr. Magn. Atmos. Elect., 50, pp. 75-78, March, 1945.—[See Abstr. 2608 (1945)].

550.389

Mean K-indices from twenty-seven magnetic observatories and preliminary international characterfigures for 1943. JOHNSTON, H. F. Terr. Magn. Atmos. Elect., 49, pp. 255-260, Dec., 1944.

550.389 2611

American magnetic character-figure, CA, three-hourrange indices, K, and mean K-indices,  $K_A$ , for July to September, 1944, and five international quiet and disturbed days for April to June, 1944. JOHNSTON, H. F. Terr. Magn. Atmos. Elect., 49, pp. 261-264, Dec., 1944.

550.389

Américan magnetic character-figure,  $C_A$ , three-hourrange indices, K, and mean K-indices,  $K_A$ , for October to December, 1944, and summary for year 1944. JOHNSTON, H. F. Terr. Magn. Atmos. Elect., 50, pp. 47-55, March, 1945.

550,389 2613

Five international quiet and disturbed days for July to September, 1944. JOHNSTON, H. F. Terr. Magn. Atmos. Elect., 50, p. 73, March, 1945.

550.822.2

Application of mud analysis logging. SOUTHER, R. E. Geophysics, 10, pp. 76-90, Jan., 1945.—The mud analysis logging system continuously analyses and records the oil and gas content of mud returns from wells being drilled by the rotary method. Oil or gas detected in the returning drilling fluid indicates oil or gas in the formation penetrated by the bit. Results of the continuous analyses are instrumentally correlated to the depths and formations from which the showings originated. A second useful phase of the system plots accurately and in detail rate of penetration or drilling speed on the log as a function of depth. Mud analysis logging can be practised in areas where the electrical log cannot be used due to high salt content or other local conditions, where dangerous hole conditions make interruption of drilling operations for coring inadvisable, and for evaluating gas zones where cores are difficult to interpret.

550.83 2615

History of the geophysical exploration of the Cameron Meadows Dome, Cameron Parish, Louisiana. McGuckin, G. M. Geophysics, 10, pp. 1-16, Jan., 1945.—The successive application, between 1926 and 1943, of various techniques to the exploration of a typical salt dome is described. These methods were: mechanical refraction seismograph (1926); torsion balance (1927); electrical refraction seismograph (1928-29); early correlation reflection seismograph (1929); dip reflection seismograph (1933); special salt profiling refraction seismograph (1942); continuous correlation reflection seismograph (1942); gravity meter (1943).

550.831

Gravimeter prospecting for chromite in Cuba. HAMMER, S., NETTLETON, L. L., AND HASTINGS, W. K. Geophysics, 10, pp. 34-49, Jan., 1945.—A report of gravimetric surveys in the chromite-bearing Camaguey District, Cuba, is given. The technical aspects of field work are discussed. Typical examples of gravity

anomalies due to chromite deposits and to various igneous rock masses are presented.

Estimating ore masses in gravity prospecting. HAMMER, S. Geophysics, 10, pp. 50-62, Jan., 1945.— The interpretation of the results of gravitational prospecting surveys is considered theoretically in terms of the magnitude of the causative mass as distinct from the conventional interpretation in terms of the mass distribution (size, shape and depth). A general proof is based on Gauss's theorem in potential theory. Practical formulae are presented for estimating the total mass directly from the gravity data, and the precision of the mass estimate is considered. The method is applied to a practical gravimetric survey over a known chromite-ore body; the estimated mass agrees with estimates from core drilling. 550.834.5

Early reflection seismograph exploration in California. SALVATORI, H. Geophysics, 10, pp. 17-33, Jan., 1945.—A brief history of the discovery of the Wilmington and Rio Bravo fields is given and the seismic maps are compared with the later geologic

maps compiled from well data.

550.837 : 621.317.39 2619

A geophysical prospecting instrument using alternating currents of audio-frequency. GUELKE, R. J. Sci. Instrum., 22, pp. 141-145, Aug., 1945.—[Abstr. 2039 B (1945)1.

550.837.3

Working depths for low-frequency electrical prospecting. Lewis, W. B. Geophysics, 10, pp. 63-75, Jan., 1945.—Electrical measurements were made on the surface of the earth with l.f. commutated current using 19 separate frequencies and 6 electrode separations. Analysis of the data indicates that there is an effect of appreciable magnitude attributable to an interface 6 000 ft, below the surface.

551.24 Contemporaneous disturbances in lacustrine beds in Kenya. KENT, P. E. Geol. Mag., Lond., 82, pp. 130-

135, May-June, 1945.

551.243.1 On the normal faulting of Rift Valley structures. DIXEY, F. Geol. Mag., Lond., 82; pp. 136-137,

May-June, 1945. 551.3.053

Erosion and water supplies. STEBBING, E. P. J. Roy. Soc. Arts, 93, pp. 290-304, May 11, 1945.— The effect of the denudation of forests and of cultivation in general upon reduction of water supplies is reviewed and illustrated by examples drawn from Britain, India, South Africa, U.S.A., Australia. The most efficacious remedy suggested against the reduction is the closure of pasturage. conserve rainfall, but it is still a matter of discussion whether they have any effect in increasing it.

551.311.234.3

J. S. G. T.

Symposium on loess. Amer. J. Sci., 243, pp. 225-303, May, 1945.—Foreword, Elias, M. K. following papers were read: Loess and its economic importance, Elias, M. K. [Abstr. 2634 (1945)]; Pleistocene loess deposits of Nebraska, Schultz,

C. B., and Stout, T. M. [Abstr. 2625 (1945)]; Glacial versus desert origin of loess, Bryan, K. [Abstr. 2626 (1945)]; A mechanical analysis of wind-blown dust compared with analyses of loess, Swineford, A., and Frye, J. C. [Abstr. 2629 (1945)]; Accumulation of dust in relation to herbaceous vegetation, Weaver, J. E.; Need of further researches on loess, Condra, G. E.; Loess types and their origin, Obruchev, V. A. [Abstr. 2627 (1945)]; Significance of loess in classification of soils, Thorp, J. [Abstr. 2633 (1945)]; Sequence of soil profiles in loess, Williams, B. H. [Abstr. 2628 (1945)]; Infiltration into loess soil, Duley, F. L. [Abstr. 2630 (1945)]; Characteristics and uses of loess in highway construction, Bollen, R. E. [Abstr. 2632 (1945)]; Observations on the properties of loess in engineering structures, Watkins, W. I. [Abstr. 2631 (1945)].

551.311.234.3 2625

Pleistocene loess deposits of Nebraska. SCHULTZ, C. B., AND STOUT, T. M. Amer. J. Sci., 243, pp. 231-244, May, 1945.—Four important loesses are recognized in the Pleistocene of Nebraska. The stratigraphy of the Pleistocene sediments is reviewed with emphasis on the placement of loess deposits in relation to marl and peat beds, buried soils, and terrace sequences. A climatic interpretation is based upon a terrace cycle concept in which loess is regarded as a product of drought and soil a result of relatively humid conditions. The Broadwater formation is described as early Pleistocene in age and consisting of a basal and upper gravel deposit separated by finer textural sediments. A loess-like silt appears in the Lisco member. A very late Pleistocene loess, the Bignell formation, is named.

551.311.234.3 2626

Glacial versus desert origin of loess. BRYAN, K. Amer. J. Sci., 243, pp. 245-248, May, 1945.—The main source of loess is the outwash of glacial rivers pulverized by frost action. The main areas of its deposition were marginal to the region most affected by the glacial born anticyclonic winds combined with prevailing westerlies. Steppe loess had its source in the adjacent deserts. The structural similarities in loesses of different origin are due to secondary processes induced by grass vegetation. Flood plains of Nebraska rivers, which in Pleistocene time were overloaded, glacial-fed streams, were the source of the loess in the State.

551.311.234.3

Loess types and their origin. OBRUCHEV, V. A. Amer. J. Sci., 243, pp. 256-262, May, 1945.—Loess is widespread in the southern part of European USSR and extends through the Kirghiz steppe to Lake Balkhash and farther north-west to Yakutia. Different types of loess are distinguished: primary loess of aeolian origin and secondary loesses, redeposited and originated by other processes. Degraded loess and compact stone-loess are recognized. Dust from which primary loess originated is believed blown by anticyclonic winds from fluvioglacial alluvium of glaciated areas and foreglacial deserts, and deposited over adjacent 'prairies. In distinction from this "cold" loess, the other, "warm" primary loess has been blown from the exposed mountain ridges and foothill thallus of Central Asiatic Mountains and adjacent

deserts, and deposited in the surrounding prairies, but part of the dust in these areas was also blown in from the distant areas of glaciation. An alternative soil-hypothesis ascribes the origin to soil-forming processes from any fine-grained formation deposited by water in foreglacial and foothill alluvial plains. Comparative petrographic, chemical, and mechanical analyses give little support to this hypothesis.

551.311.234.3

Sequence of soil profiles in loess. WILLIAMS, B. H. Amer. J. Sci., 243, pp. 271-277, May, 1945.—The time factor of soils developed from loess in western Nebraska is expressed in the character of the present soils. There is evidence that the soils on the tableland south of Chappell have developed from loess of two distinct periods of deposition.

551.311.234.3:551.311.3

2629

2630

A mechanical analysis of wind-blown dust compared with analyses of loess. SWINEFORD, A., AND FRYE, J. C. Amer. J. Sci., 243, pp. 249-255, May, 1945.— A mechanical analysis was made of wind-blown dust collected in September, 1939, from the level of the third floor of the Lakeway Hotel, Meade, Kansas. Comparisons of this analysis with previously published analyses of loess and new analyses of Kansas loess seem to demonstrate that wind can be competent to sort material to the degree represented by some loess deposits.

551.311.234.3:551.579.5

Infiltration into loess soil. Duley, F. L. Amer. J. Sci., 243, pp. 278-282, May, 1945.—Field and laboratory tests were made to determine the infiltration rate for loessial soils. When these soils are bare, they seal over during rains and have their intake rate reduced in the same way as other soils. The intake rate can be greatly increased by covering the surface with a straw mulch. The effect of the mulch was much less on the raw, parent material of the Peorian loess. The absence of a well developed structure and shortage of binding material leads to excessive erosion on this soil.

551.311.234.3:624.1 2631

Observations on the properties of loess in engineering structures. WATKINS, W. I. Amer. J. Sci., 243, pp. 294-303, May, 1945.—Some personal observations on the properties of loess and loess-like materials which affect their behaviour in engineering structures. Most important of these are texture, chemical composition and the amount of moisture absorbed and held.

551,311,234,3:625.8

Characteristics and uses of loess in highway construction. Bollen, R. E. Amer. J. Sci., 243, pp. 283-293, May, 1945.—Some of the properties important in highway construction are described and discussed. Special physical tests of soils, which include loess, and which have been standardized by engineers, permit identification of their types and their expected performance as building material. Disturbances of soils due to exposure in cuts and to various subsequent engineering operations have important effect on their properties. Performance of loess in subgrades, as binder in stabilized base courses and filler in bituminous surfacing are described.

· 2637

551.311.234.3:631.4

2633

Significance of loess in classification of soils. THORP, J. Amer. J. Sci., 243, pp. 263-270, May, 1945.—The soils developed from loess vary considerably because of great variation in combination of factors involved in soil formation: climate, biological activity, relief and time. Besides, the character of loess, which is the parent material, also varies considerably. Drawing of the boundaries of types of soils in the soil survey of the Great Plains is helped considerably by the understanding of loess origin, thickness, and distribution.

551.311.234.3:631.4:625.8 2634

Loess and its economic importance. ELIAS, M. K. Amer. J. Sci., 243, pp. 227-230, May, 1945.—Fertility of loess is its most valuable property. The principal wheat producing regions of the world roughly coincide with the areas of distribution of this golden earth of agriculture. Where it mantles the ground, the loess and the soils developed from it are the principal or, frequently, the only readily available material for roads, dams, and other engineering constructions.

551.311.3: 551.311.234.3 see Abstr. 2629

551.463: 535.345.1 see Abstr. 2245

551.510.535: 523.74: 537.311.37 see Abstr. 2301

551.574: 536.42 see Abstr. 2273

551.579.5 : 551.311.234.3 see Abstr. 2630

553.621 : 535.375.5 see Abstr. 2251 553.621 : 539.32 see Abstr. 2404

553,621: 548.24 see Abstr. 2573

576.858: 582.951.4: 621.384.4: 535.212

see Abstr. 2216

576.858.8: 548.73 see Abstr. 2593

577.15: 541.124 see Abstr. 2450

578.088.9: 534.321.9 see Abstr. 2206

581.036 : 541.144.7 see Abstr. 2490

582.951.4: 576.858: 621.384.4: 535.212 see Abstr. 2216

591.1.044: 539.166.92 see Abstr. 2377

591.112: 532.57 see Abstr. 2178

591.18 2635

Neural mechanisms for the concepts of difference and similarity. LANDAHL, H. D. Bull. Math. Biophys., 7, pp. 83-88, June, 1945.—A mechanism is described which reacts to differences in different modalities of a stimulus pattern. Another mechanism reacts to similarity; this is measured by the number of neurons common to the 2 stimulus patterns.

L. S. G. 591.18

A heterarchy of values determined by the topology of nervous nets. McCulloch, W. S. Bull. Math. Biophys., 7, pp. 89-93, June, 1945.—The function of reflexes, i.e. activities with a feed-back mechanism, includes all purposive activities. The latter are of dromic character and so the closed circuits sustaining them and their interaction may be treated topologically. The value anomaly is studied where A is preferred to B, B to C, but C to A. The apparent inconsistency of preference may be submitted to finite tolological analysis based on the finite number of nervous cells and their possible connections. L. S. G.

591.181:537.39

Effects of constant current in relation to "functional polarity." WINTERSTEIN, H., AND SEDEFCIYAN, A. Nature, Lond., 155, pp. 238-239, Feb. 24, 1945.—Scheminzky and his co-workers found that a descending (head-to-foot direction) constant current passed through a frog produced paralysis (galvano-narcosis) and an ascending current produced convulsions. He attempted (Pfügers Arch., 243, 439-456, 1940) to explain this by a "functional polarity," but experi-

ments here recorded contradict and refute this. C. J. G. 611.018 2638

A note on the physiological arrangement of tissues. MORALES, M. F., AND SMITH, R. E. Bull. Math. Biophys., 7, pp. 47-51, June, 1945.—A presentation of the kinetics of uptake of an inert solute by tissues arranged in series, distinct parallel and competitive parallel, with respect to the circulation. The uptake is described by a series of exponentials, the number of terms and the constants depending upon the arrangement. The degree of coupling is reflected in the nature of the constants.

L. S. G.

611.7 2639

Dynamics of quadrupedal locomotion. HOUSE-HOLDER, A. S. Bull. Math. Biophys., 7, pp. 53-57, June, 1945.—Fischer's dynamical equations for unbranched systems in two dimensions are extended to the branched system exhibited by a quadruped.

L. S. G.

612.76: 531.1 see Abstr. 2147

612.84 2640

A theory of photosensitivity of some lower animals. Runge, R. Bull. Math. Biophys., 7, pp. 59-67, June, 1945.—The two-factor nerve excitation theory is used with the assumption that the threshold itself varies under the action of light, thus providing a third factor. A study is made of the dependence of the reaction time upon (i) the intensity of the stimulus, (ii) time of exposure to light of a fixed intensity, and (iii) dark adaptation. Light adaptation and the effects of termination of exposure after light adaptation are discussed. The theory is in agreement with observations.

L. S. G.

612.84: 535.7 see Abstr. 2262

612.84: 535.733.1 see Abstr. 2265

613.6: 535.7 see Abstr. 2263

614.841 : 542.978 see Abstr. 2535

314.041 . 342.910 see Ausii. 2333

Outline of a mathematical approach to the cancer problem. RASHEVSKY, N. Bull. Math. Biophys., 7, pp. 69-81, June, 1945.—A relationship is derived which gives the probability of cancer incidence as a function of age. The relation is in agreement with the rather inadequate empirical data. A relation for the speed of wound healing as a function of age is also derived and is in agreement with observational data. Possibilities of practical applications are discussed.

L. S. G.

616.9: 541.127.2 see Abstr. 2462

620,193,19 2642

The moisture content distribution in wood used as a partition between water and air. ROBERTS, H. D. J. Coun. Sci. Industr. Res. Aust., 17, pp. 133-138, Aug., 1944.—Six timber species were used in these

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experiments, the results of which indicate that when one face of a 2 in, wooden partition is kept continuously wetted, and the other face continuously dry (such as occurs in vats, casks, tanks, wood pipe, ship timbers etc.) a very high moisture gradient exists across the thickness of the partition once equilibrium is attained. In the species tested the average moisture content of the 0.2 in, thickness of wood nearest the wetted face tended to reach the average green moisture content of the species, whereas the average moisture content of the 0.2 in. thickness nearest the dry face (in contact with air) tended towards an average moisture content of about 14%. At a distance from 0.5 to 0.75 in. from the wetted face, the moisture content tended to reach equilibrium at approx. the fibre saturation point for the species used. C. J. G.

620.193.92 : 541.132.3 see Abstr. 2476

624.15: 531.2 see Abstr. 2148

631.4:551.311.234.3 see Abstr. 2633

631.4:625.8:551.311.234.3 see Abstr. 2634

631.42

The use of the hydrometer for the mechanical analysis of soils. Downes, R. G. J. Coun. Sci. Industr. Res. Aust., 17, pp. 197-206, Aug., 1944.-The theory of hydrometers is discussed and the various possible sources of error considered. These are: (i) those associated with the relation of density to percentage of soil in suspension; (ii) those associated with the determination of the depth in the suspension at which the density is the same as that measured by the hydrometer; (iii) those due to practical difficulties. Results from a test hydrometer designed to minimize these errors have been compared with results obtained by the standard pipette technique for a number of soils. The mean of all differences expressed as a percentage of the hydrometer reading and irrespective of sign, is about 5%. This may be considered a creditable performance when allowance is made for the differences in dispersion technique used for the two methods. C. J. G.

632.94: 541.42 see Abstr. 2510

661.713: 532.13 see Abstr. 2170, 2171

662.41 : 541.126 see Abstr. 2455

662.66: 539.501 see Abstr. 2429

663.63: 541.132.3 see Abstr. 2477

Industrial uses and marketing problems of tall oil. MCKEE, R. H. Paper Tr. J., 120, TAPPI Sect., pp. 35-36, Jan. 25, 1945.—Industrial applications include uses in the soap, varnish, paint and textile industries, in cutting oils, lubricants, insect spray, asphalt and disinfectant emulsifiers, and in oils and plastics. Tall oil has now been shown to be a mixture of linolenic and linoleic acids with a small amount of oleic acid, and a non-acidic portion containing

2,2-dihydrostigmasterol and lignoceryl alcohol. The

proportions of these vary with the wood, cooking conditions and refining methods used.

J. G.

665.52 : 542.48 : 545.727 see Abstr. 2545 666.1 : 532.13 : 539.32 see Abstr. 2403

666.1 : 539.56 see Abstr. 2434

666.1: 541.132 see Abstr. 2473

666.1.038: 539.319 see Abstr. 2401

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667.4

7.4 2645 Modified pen-and-ink sizing test. SMITH, B. W.

Paper Tr. J., 120, TAPPI Sect., pp. 72-73, Feb. 22, 1945.—Tests for sizing (ink feathering) made with a standard ink may be very unreliable, inasmuch as modern inks vary considerably in their capacity to produce feathering. It is advisable to use (in addition to the TAPPI Standard T431 m black ink) a red, green and violet aq. ink free from and containing 0.25, 0.5, 0.75 and 1.0% of a surface active material (Na<sub>3</sub>PO<sub>4</sub>, PhOH, or diethylene glycol, respectively). Paper which does not feather with any of these inks containing 0.5% of surface-active reagent is completely satisfactory. The results are independent of the pen point if a ball-pointed nib is used, but each mark should have the form of a heavy right-angled X, each arm of which is at 45° to the paper grain. J. G.

667.7: 539.574 see Abstr. 2435

669.018: 532.782 see Abstr. 2193

669.018: 539.43 see Abstr. 2420

669.14: 539.62 see Abstr. 2438

669.14: 544: 535.247.4 see Abstr. 2226

669.14: 548.526: 539.378.3 see Abstr. 2405

669.215 : 541.122.2 see Abstr. 2440

669.721 : 539.4.016.3 see Abstr. 2418

669.721.5: 548.73 see Abstr. 2594

674.03: 539.384.4 see Abstr. 2409

676 : 620.115

New apparatus for determining the water vapour permeability of sheet materials. Cornwell, R. T. K. Paper Tr. J., 120, TAPPI Sect., pp. 82-93, March 1, 1945.—[Abstr. 1811 B (1945)]:

(76.1 541.104 4/ 4 0451

676.1 : 541.124 see Abstr. 2451 676.2

2647

St. Regis successfully bleaches groundwood. LEE, J. A. Chem. Metall. Engng, 51, pp. 106-108, 1944.— The use of existing equipment in a newsprint paper mill for bleaching groundwood by the peroxide process, is described. The resulting paper is superior to newsprint made from ordinary groundwood in colour, cleanliness and strength, slightly inferior to it in opacity, and equal to it in other respects. The reagent used contains Na<sub>2</sub>O<sub>2</sub> (1-2% on wt. of dry pulp), with Na<sub>2</sub>SiO<sub>3</sub> (as detergent, penetrant, stabilizer, and corrosion preventive), and MgSO<sub>4</sub> (to prevent catalytic decomposition of the peroxide by the metals present).

676.3 2648

Use of pyrodextrins in the clay coating of paper, KERR, R. W., AND SCHINK, N. F. Paper Tr. J., 120, TAPPI Sect., pp. 77-80, Feb. 22, 1945.—The colloidal characteristics of amylopectin are such as to make it more desirable than is amylose for use in clay coatings. Dextrinization involves the linkage of linear chains (from amylose) with branched structures (from amylopectin) to form a pyrodextrin, which has coating qualities superior to either. Thus, pyrodextrins made from corn starch produce clay mixes of low viscosity and plasticity, but high adhesive value, and they are suitable for both low- and high-density coating machines. Advantages of the latter are rapidity of drying, high machine speeds, and a high finish on the final paper.

676.3\*: 535.683.1

2649

Standard testing methods for light fastness in the paper industry. LIPS, H. A. Paper Tr. J., 120, TAPPI Sect., pp. 108-110, March 15, 1945.—Previous work is reviewed. As a method of standardizing fading lamps, it is proposed that a piece of paper or cloth (the control), suitably dyed, should be exposed to the lamp for 20 hr. and then matched against one of a set of permanently coloured plastic chips which will illustrate the results of 12, 14 etc. to 26 hr. of exposure in a standard lamp. If therefore the control is exposed with the sample for 20 hr. in the lamp to be standardized and again matched against the chips, the exposure time can be reported in terms of the corresponding no. of standard hr. The method may be used to rate exposures to sunlight, and to relate such exposures to fading-lamp tests.

676.3: 539.536 see Abstr. 2432

2650 676.6

Development of weatherproof, solid fibre board. BARKER, H. T. Paper Tr. J., 120, TAPPI Sect., pp. 117-119, March 22, 1945,-A brief survey of the development and future possibilities of these boards. with special reference to specifications (caliper, dry and wet bursting strengths, and splitting resistance) for V1, V2 and V3 grades. Asphalted duplex liners provide the cheapest and simplest means of producing high wet strength without loss of rigidity in the wet state. For the measurement of H2O resistance the Cobb and immersion tests are preferred. Polyvinyl alcohol base adhesives are used for laminating weatherproof boards. Almost as satisfactory, but requiring closer control in use, though much cheaper, is an acid starch which is heated at 90-91°, cooled, and mixed with 5-20% of U-F resin.

677: 519.283: 539.216.1 see Abstr. 2386

677: 539.217.1 see Abstr. 2388

677.01: 539.217.3: 532.696.1 see Abstr. 2187

677.2: 539.42: 539.216.1 see Abstr. 2385

677.3: 519.21 see Abstr. 2100

677.65: 539.217.5 see Abstr. 2389

678: 539.388.8 see Abstr. 2412

678: 539.431 see Abstr. 2422

678: 539,501 see Abstr. 2430

678.7: 539.388.8 see Abstr. 2413

678.7: 548.73 see Abstr. 2595

679.5 = 3

The formation of synthetic resins from Schiff's bases. I. Resins from aromatic sulphonic acid amines and formaldehyde. II. Further investigations on the nature of sulphonamide resins. Scheele, W., and Steinke, L.

Kolloid Z., 97, pp. 176-189, Nov., 1941, and 100, pp. 361-368, Sept., 1942.

679.5: 539.217.2: 539.216.2 see Abstr. 2387

679.5: 539.4.01 see Abstr. 2417

679.5: 539.434 see Abstr. 2423

679.5: 621.383.4: 539.385.08 see Abstr. 2411

2652 Vulcanization of saturated acrylic resins. MAST,

W. C., REHBERG, C. E., DIETZ, T. J., AND FISHER, C. H. Industr. Engng. Chem., 36, pp. 1022-1027, Nov., 1944. 679.56

Vulcanized acrylic resins. Copolymers of ethyl acrylate and allyl maleate. MAST, W. C., SMITH, L. T., AND FISHER, C. H. Industr. Engng Chem., 36, pp. 1027-1031, Nov., 1944.

681.4:535.87 see Abstr. 2269

770.1: 541.141 see Abstr. 2486

771.35: 535.317 see Abstr. 2230

771.35: 535.81 see Abstr. 2266

771.37:621.383

2654

Direct-reading colour densitometer. Sweet, M. H. Electronics, 18, pp. 102-106, March, 1945.-[Abstr. 2135 B (1945)],

778:535.33-31

2655 Spectrographic plates of substantially uniform contrast in the ultra-violet. DAVEY, E. P., AND GAUNT-LETT, M. D. J. Soc. Chem. Ind., 64, pp. 70-72, March, 1945.—It is shown that very strong light absorption by the silver halide in the region 2 500-3 100 Å confines the photographic image largely to the surface of the plate, and that the usual increase in contrast above this wavelength results from a lesser absorption and a greater penetration of the light into the emulsion layer. By restricting the image to the surface here also, while at the same time not decreasing the contrast at lower wavelengths, plates of substantially uniform contrast have been prepared for use in spectrography, and the results of measurements on such a plate are given.

778.3:621.386.84

2656

Millionth-of-a-second X-ray snapshots. C. M., EHRKE, L. F., AND ZAVALES, C. T. Westingho. Engr, 5, pp. 99-103, July, 1945.—[Abstr. 2169 B (1945)].

778.33:621.383:621.396.664

2657

Electronic X-ray timer. Moreland, H. D. Radio-Electronic Engng, 4, pp. 14-15 and 45-46, June, 1945.— [Abstr. 2221 B (1945)].



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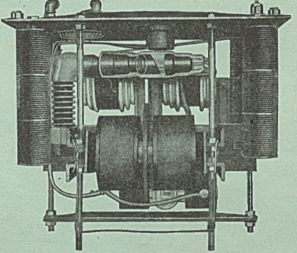
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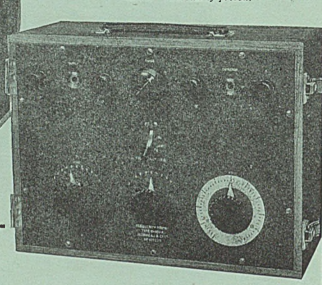
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