Annotated Accession
List of Data Compilations
of the NBS Office of
Standard Reference Data
TECHNICAL NOTE 554

ISSUED SEPTEMBER 1970

CODEN: NBTNA

Annotated Accession List of Data Compilations of the Office of Standard Reference Data

Herman M. Weisman and Gertrude B. Sherwood

Office of Standard Reference Data
National Bureau of Standards
Washington, D.C. 20234

NBS Technical Notes are designed to supplement the Bureau’s regular publications program. They provide a means for making available scientific data that are of transient or limited interest. Technical Notes may be listed or referred to in the open literature.
ABSTRACT

The National Bureau of Standards, Office of Standard Reference Data has attempted to acquire all significant reference data compilations on a worldwide basis. This publication lists the collection of documents so acquired, together with their abstracts. The documents are organized in the following categories: General Collections, Nuclear Properties (including Fundamental Particles Properties), Atomic and Molecular Properties, Solid State Properties, Chemical Kinetics, Colloid and Surface Properties, Mechanical Properties, and Thermodynamic and Transport Properties. Sources of availability for the listed publications are also provided.

Key words: Accession List, Atomic and Molecular Properties, Chemical Kinetics, Colloid and Surface Properties, Fundamental Particles Properties, General Collections, Mechanical Properties, Nuclear Properties, Solid State Properties, Thermodynamic and Transport Properties
## Table of Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>1</td>
</tr>
<tr>
<td>General Collections</td>
<td>4</td>
</tr>
<tr>
<td>Nuclear Properties (Including Fundamental Particles Properties)</td>
<td>18</td>
</tr>
<tr>
<td>Atomic and Molecular Properties</td>
<td>57</td>
</tr>
<tr>
<td>Solid State Properties</td>
<td>94</td>
</tr>
<tr>
<td>Chemical Kinetics</td>
<td>121</td>
</tr>
<tr>
<td>Colloid and Surface Properties</td>
<td>132</td>
</tr>
<tr>
<td>Mechanical Properties</td>
<td>135</td>
</tr>
<tr>
<td>Thermodynamic and Transport Properties</td>
<td>141</td>
</tr>
</tbody>
</table>
ANOTATED ACCESSION LIST
of DATA COMPILATIONS of the
NBS Office of Standard Reference Data

Introduction

The scientist or engineer seeking specific data is often confronted with a number of difficulties. Where can the data be found? Are they complete? Are they compatible with other sources? Most important, how reliable are they? Each year the problem for the scientist and engineer becomes greater. As the volume of literature expands, he finds even more difficulty in trying to keep abreast of literature relevant to his field. This is why the National Standard Reference Data System (NSRDS) was created. The NSRDS is a nationwide program to help provide the U.S. technical community optimum access to the quantitative data of the physical sciences, critically evaluated and compiled in convenient formats. It was established in 1963 by action of the President's Office of Science and Technology with responsibility for administering the effort given to the National Bureau of Standards. The Office of Standard Reference Data was set up within NBS to manage and coordinate the program. The National Standard Reference Data System operates, for the most part, as a decentralized cooperative network of data centers and data compilation projects for which the Office of Standard Reference Data (OSRD) is the program management vehicle. The data centers are composed of individuals or groups working on the collection and evaluation of all the data within specific technical areas. These centers, many of which are located in the experimental divisions of the NBS, some in other government laboratories, at universities or at private companies, are the source of the critically evaluated compilations, reviews, and bibliographies published by the NSRDS program.

The scope of the program has been limited to the physical sciences. For management purposes, the Office of Standard Reference Data has divided its activity into specific technical areas: nuclear data, atomic and molecular data, solid state data, thermodynamic and transport data, chemical kinetics, mechanical properties, and colloid and surface properties. Each of the technical areas is organized in a similar way, having a program manager who monitors data compiling groups who may be working at the NBS or may be under contract, or who might be a group sponsored by the other government agencies. Each area program has an advisory panel of expert consultants, who aid in appraising activities, assessing priorities and recommending what the NSRDS should do in a given area.

Among the primary duties of the Office of Standard Reference Data, according to the directive from the Office of Science and Technology which established the program is the operation of an information services center for the NSRDS. Because the major available resources of the NSRDS program have been applied thus far to the production of compilations and critical reviews, the development of a vigorous information service activity has been delayed. Nevertheless, some basic and significant services have been implemented. One of the more active is the operation of a Data File. The mission of the OSRD Data File (library) is to acquire selectively, on a world-wide basis, reference data compilations and to organize the collection systematically. The Data File houses a collection of about 1300 reference data compilations, critical reviews, bibliographies and other ancillary reference works. It has been judged as one of the most extensive collections of its kind in the world.

Systematic compilations of data contribute in a fundamental way to the progress of science and technology. Data Compilations have been among the
basic tools of scientists and engineers throughout the history of technology. Each worker owns at least one handbook containing, among other useful information, table after table of data on the properties of the substances and systems that he deals with daily. Recognizing that knowledge of the items within its collection could be of value to many scientists and engineers, the Office of Standard Reference Data has organized the documents and prepared an annotated accession list to be made available to the technical community. Individuals as well as organizations might be interested in single items within the collection of groupings of them.

The Annotated Accession List has been organized in accordance with the present shelving arrangements of the documents in the Office of Standard Reference Data library. We consider this a first approach to classification and indexing of the holdings of the program, while searching for a classification and indexing system that can withstand the test of file-growth, time, and complexity. A modest examination has shown that no existing classification scheme is satisfactory for NSRDS needs. The present size of the Data File and its highly specialized nature and use were important factors in the evolvement of the scheme presently being used. Classifying publications for shelf location and for convenient browsing is a somewhat different problem than that of indexing contents to permit rapid location of individual data points.

A classification scheme for shelving does not need to have a large number of partitions of a subject matter. Factors in the development of the system included the needs of the user, his familiarity with or capability to become acquainted with the shelving system, and his subjective reactions while browsing. For these reasons, the approach to shelving the holdings of the Office of Standard Reference Data File has emphasized the needs of those who use the file for browsing -- that is, the professional staff of the Office.

Nine classification categories based on the NSRDS technical areas are used. The categories are further partitioned by a decimal type of breakdown. The scheme is as follows:

- General Collections
- Fundamental Particles Properties
- Nuclear Properties
- Atomic and Molecular Properties
- Solid State Properties
- Chemical Kinetics
- Colloid and Surface Properties
- Mechanical Properties
- Thermodynamic and Transport Properties

Associated with the classification numbers are two additional digits. These last two digits are shelving control codes. With the exception of the General Collections, Fundamental Particles Properties and Nuclear Properties categories, the number after the decimal point indicates the type of compounds discussed in the publication: .1 indicates only inorganic compounds, .2 both inorganic and organic, and .3 only organic compounds. For example, the classification 3.1 indicates that the publication deals with atomic and molecular properties of only inorganic compounds, 8.2 indicates the publication contains information on thermodynamic and transport properties of both inorganic and organic compounds.

The Fundamental Particles Properties and Nuclear Properties are subdivided by the categories: stationary states (1.1 and 2.1) and processes (1.2 and 2.2). Further, nuclear properties, stationary states, and processes are subdivided by a particular state (i.e., quantum numbers, 2.15) or a particular reaction (i.e., interactions with neutrons, 2.25).
The General Collections classification contains publications appropriate to several categories, housing material like data collections (0.11), handbooks (0.21), and bibliographies (0.31).

The classification number is preceded by the document's accession number and appears on the first line of the item's listing. The items in the Annotated Accession List are arranged by the nine subject categories in accordance with the indicated scheme. For the convenience of the reader, items shelved in one technical area but applicable to still another or several have been repeated in the separate area listings.

This Annotated Accession List does not include an index. Its classification scheme is intended essentially as an aid to shelf retrieval of documents. However, the decimal breakdown used provides limited capability for substance-property retrieval. The Office of Standard Reference Data is, at present, beginning exploratory work on substance-property retrieval.

We have attempted to provide information on the sources of availability for the listed publications. To conserve space, the Clearinghouse for Federal Scientific and Technical Information (CFSTI), the Defense Documentation Center (DDC), the Superintendent of Documents, Government Printing Office (GPO), and the Division of Technical Information, Atomic Energy Commission, (DTI, AEC) have been abbreviated as indicated.

We wish to emphasize that the Office of Standard Reference Data does not necessarily vouch for the validity, criticality, or currency of the data within these documents. They have been selected on the basis of the judgment by the technical staff of the Office as providing some information or data of a useful and reference nature. Most of the annotations are based on author abstracts in the original documents.

The Office of Standard Reference Data has attempted to acquire all significant reference data compilations in existence on a world-wide basis. Availability and financial considerations may have precluded acquisition of some documents which properly might have been included. The Office of Standard Reference Data invites readers to inform it of any additions which might enhance the collection.

The compilers of this Annotated Listing want to acknowledge their indebtedness to a number of individuals without whose help this Listing would not have been possible. These individuals include: Edmonia Kearney, Lyle D. Joyce, Allen M. Weinberger, Alfred Weissberg, David R. Lide, Jr., Edward L. Brady, Howard J. White, Jr., L. H. Gevantman, David T. Goldman, and Joseph Hilsenrath. We want to express our appreciation to Robert Thompson, Carla Messina, and Robert McClennon for the help in mechanizing the Annotated Listing. A special note of thanks is due to Mrs. Darlene Graham for keyboarding the items in this List and to Mrs. Roberta Kadesch, Mrs. Barbara Sommervill, Leonard Loewentritt and Elizabeth Wilson for the painstaking and diligent proofreading.
General Collections
This Handbook contains tabulations of mathematical methods, mathematical functions, and tabulations of their values. The explanatory sections were written by various authors; the tables were calculated by means of a digital computer. The following functions have been tabulated: logarithmic, exponential, trigonometric, hyperbolic, exponential integral, gamma function, Legendre functions, Bessel functions of integer and fractional order, integrals of Bessel functions, Struve functions, confluent hypergeometric functions, Coulomb wave functions, hypergeometric functions, Jacobian elliptic functions, theta functions, elliptic integrals, Weberstrass elliptic and related functions, parabolic cylinder functions, Mathieu functions, spheroidal wave functions, orthogonal polynomials, Bernoulli and Euler polynomials, Riemann zeta function. The material analyzes, numerical interpolation, differentiation and integration, probability functions, miscellaneous functions, scales of notation and Laplace transforms.

This book presents recommended values for astronomical and astrophysical quantities. These values were arrived at from an examination of literature published through 1955. The following types of data have been included: general constants and units, atoms, spectra, radiation, the earth, planets and satellites, interplanetary matter, the sun, normal stars, special stars, stellar populations, interstellar space, star clusters, galaxies, and groups of galaxies.

This is a preliminary and academic exploration of approaches to information handling by the Office of Standard Reference Data. It speculates on means for the selection, acquisition, intellectual organization, and storage of the information which might underlie the Information Services Operation of the National Standard Reference Data System, as well as for methods of locating desired information items in storage, retrieving, and displaying or communicating them. Questions of the use of computers for these purposes are discussed, including selection of equipment, arrangement of digital storage, input format, remote access, and the economics of choosing certain functions of the system for mechanization. Also an interim system based on conventional and, in the main, manually operated files, is described.

These loose-leaf documents, the AIChE Physical Property Estimation System, are manuals directed primarily at computer oriented staff who are familiar with the operation of the computer system on which programs are compiled and executed. Volume II instructs the user on how he can communicate with the estimation system. The input requirements, alternatives, and options are described together with the various outputs that the system can generate. Volume II further describes how the system programs are delivered and how they can be compiled, assembled into a system, and executed on a computer. Volume III discusses the process of updating the system, system table requirements, the subject of transfer function linking, and the processing of information from program to program through a pass tape system. This volume contains a functional description and flow charts for the main system programs. The layout of common storage for each main program is also shown. Chapter 8 discusses the reliable techniques to estimate surface tensions of pure liquids, the variation of surface tension with temperature and pressure, and surface tension estimation methods for non-aqueous mixtures and aqueous mixtures. Interfacial tensions are briefly considered. Chapter 9 deals with the viscosity of gases and liquids. Methods are recommended for correlating low-pressure viscosity data with temperature estimating viscosities when no experimental data are available, estimating the effect of pressure on viscosity, and estimating the viscosities of mixtures. The molecular theory of viscosity is also considered.
This report summarizes the status of activities of the NBS Office of Standard Reference Data as of April 1968. It provides a detailed review of the data compilation activities within the seven broad categories of the technical scope of the program: (1) nuclear data, (2) atomic and molecular data, (3) solid state data, (4) thermodynamic and transport data, (5) chemical kinetics, (6) colloid and surface properties, and (7) mechanical properties. Progress in data systems design and development and in information services are reviewed. Certain problem areas of the program are identified. The appendix includes: (1) a listing of information and data centers associated with the Office of Standard Reference Data, (2) publications of the National Standard Reference Data System, and (3) a listing of organizations, groups, or individuals compiling or evaluating data.

In June 1963, the National Bureau of Standards was assigned the responsibility, by the President's Office of Science and Technology, of administering a government-wide program to promote and coordinate systematic data compilation and evaluation activities in all fields of the physical sciences. This government-wide program has been called the National Standard Reference Data System. The present report describes the background of this program and the mode of approach adopted by the management of the National Bureau of Standards in fulfilling its responsibility.

This Handbook contains data on the properties and uses of materials, the elements of material economics, and miscellaneous topics. The materials properties and uses section makes up the largest section of the book, including physical, chemical, and mechanical properties, as well as uses of the materials. A materials index is provided.

A bibliography of 2,482 references to published research on the properties of deuterium and tritium compounds is given. The subject matter of each entry in the bibliography is indicated by letter codes related to a list of broad subject headings shown in the Introduction. An index of deuterium and tritium compounds and a subject index are included.

The Handbook of Physics is a compendium of articles and tabulations of data, written by about 90 contributors, on many topics of physics. Among the topics considered are: mathematics, mechanics of particles and rigid bodies, mechanics of deformable bodies, electricity and magnetism, heat and thermodynamics, optics, atomic physics, solid-state physics, and nuclear physics. Tables and graphs of data are included.

The Consolidated Index is a key to the contents of publications that present critically evaluated numerical property values. This initial volume contains in codified form the contents of the following six compilations of physical and thermodynamic data: Selected Values of Properties of Hydrocarbons and Related Compounds (American Petroleum Institute Research Project 44), Selected Values of Properties of Chemical Compounds (Manufacturing Chemists Association Research Project), Selected Values of Chemical Thermodynamic Properties (NBS Circular 500), Thermodynamic Properties of the Elements (D.R. Stull and G.C. Sinke), Contributions to the Data on Theoretical Metallurgy (Bureau of Mines Bulletins 383, 384, 395, 406, 407, 477, 542, 584), and Selected Values for the Thermodynamic Properties of Metals and Alloys (Minerals Research Laboratory, University of California). Each element, compound, or system of pure substances appears as a separate index entry. For a given substance, all properties that appear in one or more of the compilations are listed separately under each. A given property, if appearing in several, is repeated under each publication.

This is a preliminary report to the National Research Council consisting of a reevaluation of the atomic constants and conversion factors of physics. As a result of the precision microwave and atomic beam techniques, a great advance has been accomplished in the accuracy of such constants as the charge and mass of the electron, h, Planck's constant, Avogadro's number, the velocity of light, the conversion factor from the Siegbahn nominal scale of x-units to milliangstroms, and a host of some sixty or more useful derived constants and conversion factors which can be computed from the preceding with the use of accurately known other constants, such as the Rydberg, the gas constants, etc. In this report the least-squares method was employed to adjust the data to arrive at
In this book the authors summarize the many experimental observations on water and present theoretical discussions which effectively correlate the data. The topics covered include the water molecule and forces between water molecules; the thermodynamic properties of steam; the structures of the ices; the thermodynamic, electrical, spectroscopic and transport properties of the ices and of liquid water; hydrogen bonding in ice and water; and models for liquid water. The authors emphasize the relation of the properties of ice and water to their structures, and discuss properties such as infrared and Raman spectra because they reveal a great deal about these structures. The inclusion of some background material in physical chemistry makes the book accessible to the biologist, biochemist, and geologist as well as the chemist and physicist. A bibliography listing a number of recent articles on the structures and properties of water completes the volume.

The Smithsonian Physical Tables consist of 901 tables giving data of general interest to scientists and engineers. The topics covered in the tables include general physical constants, units of measurement, photometry, characteristics of light sources, emissivity, densities, boiling and freezing points of materials, specific and latent heats, mechanical properties of materials, viscosity, electrical properties of materials, dielectric properties of materials, magnetic properties of materials, optical properties of materials, air and the atmosphere, kinetic theory of gases, atomic and 10-9 molecular physics, radioactivity, X-rays, fission, cosmic rays, gravitation, temperature, solar radiation, aviation, astrophysics, and oceanography.

These tables list the melting and boiling points and indices of refraction of organic compounds. The tables are arranged by class of compound, and by melting and boiling point within each class. The classes of compounds considered are: alkanes and cycloalkanes, alkenes, cycloalkenes and dienes, alkyynes, aromatic hydrocarbons, halides, alcohols, phenols, others, aldehydes, ketones, quinones, carboxylic acids, carboxylic acid halides, anhydrides, amides, imides, esters, amino acids, amines, carbohydrates, nitrile compounds, nitriles, and sulfuric acids.
density of aqueous solutions, dipole moments, heat capacity of solids, latent heats and thermodynamic functions. The atomic physics data includes ion and electron mobilities, recombinations and diffusion coefficients, electron configurations of elements, ionization potentials, X-ray spectra, and atomic constants. The mathematical functions tabulated include: logarithms, antilogarithms and exponential functions.

#6604006 *0.2143
Kitt, B.; Evered, D. S.
ROCKET PROPPELLANT HANDBOOK
New York: The Macmillan Company
1960
334p
This book contains data on the chemical propellants used in rocket engines. It also includes a general discussion of rocket propulsion and of liquid and solid propellants. The following inorganic propellants are discussed: aluminum, ammonia, boron compounds, fluorine and related compounds, hydrazine, hydrogen, hydrogen peroxide, lithium, lithium hydride, nitrates, nitric acid, nitrogen oxides, oxygen and ozone, perchlorates and water. The following organic compounds are discussed: alcohols, ethers, amines, benzene, metal-organic compounds, nitroparaffins, nitrates, petroleum fractions, polymers, saturated hydrocarbons, and unsaturated hydrocarbons. The properties tabulated for each compound include none or all of the following: molecular weight, color, odor, melting and boiling point, critical point, triple point, coefficient of thermal expansion, vapor pressure, surface tension, heats of formation and vaporization, heat capacity, heats of formation and combustion, heat of solution in water, free energy of formation, entropy, enthalpy, free energy, viscosity, thermal conductivity, velocity of sound, magnetic susceptibility, electrical conductivity, dielectric constant, dipole moment and index of refraction.

#6508013 *0.2143
Knowlton, A. E. (Editor)
STANDARD HANDBOOK FOR ELECTRICAL ENGINEERS, NINTH EDITION
1957
2230p
This handbook contains data useful in the field of electrical engineering. The topics covered by the handbook are: units and conversion factors, electric and magnetic circuits, measurements, properties of materials, circuit elements, transformers, regulators, reactors, alternating current generators and motors, direct-current generators and motors, rectifiers and converters, power-plants, power economics, power system, electrical equipment, power transmission, power distribution, wiring design for commercial and industrial buildings, illumination industrial and commercial power, electric heating and welding, electricity in transportation, electrochemistry and electrolytic cells, batteries, wire telephone and telegraph, electronics and electron tubes, radio, radar and carrier communication, codes and standard practices, and electro-physics.

#6800005 *0.3241
Kuchowicz, B.
NUCLEAR ASTRONOMY A BIBLIOGRAPHICAL SURVEY
New York: Gordon and Breach, Science Publishers
1967
450p
A survey is given on the abundance of chemical elements, nuclear reactions in stars and stellar energy sources, stellar structure and evolution, solar system, nova and supernova, x-ray and stellar matter, nucleosynthesis, neutrino emission by stars, origin of cosmic radiation, radio and gamma astronomy, cosmology and cosmogony problems, and related nuclear physics. Twenty-one hundred and one papers are quoted from 1335 to the present.

#6500525 *0.1151-3.151
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME I: ATOMIC AND MOLECULAR PHYSICS, PART 1: ATOMIC AND IONS
Eucken, A.; Hellewege, K.-H. (Editors)
Berlin, New York: Springer-Verlag
1950
441p
In German
This section of the Landolt-Bornstein tables contains data on the fundamental physical constants and the properties of atoms and ions. Information for the use of the tables, including data on systems of measurement and units, has also been included. A discussion of atomic spectra, including wavelength standards, electron terms, important spectral line wavelengths, ionization potentials, electron affinities, X-ray spectra, Zeeman and Stark effect data, pressure broadening and pressure displacement of spectral lines, and oscillator strengths and lifetimes of excited states is presented. Other properties of atoms and ions compiled include electron distribution, X-ray scattering and absorption, cross sections, magnetic moments, magnetic electric polarizability, and Faraday effect data.

#6500526 *0.1151-3.151-5.151-8.151
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME I: ATOMIC AND MOLECULAR PHYSICS, PART 2: MOLECULES I (NUCLEAR STRUCTURE)
Eucken, A.; Hellewege, K.-H. (Editors)
Berlin, New York: Springer-Verlag
1951
571p
In German
This section of the Landolt-Bornstein tables contains atomic physical data of molecules which describes the structure and dynamics of their nuclear framework. The following properties have been tabulated: atomic distances and structures, valence energies of chemical bonds, dissociation energies of chemical bonds, and molecular vibrations and rotations.

#6500527 *0.1151-2.2051-3.151
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME 3: MOLECULES II (ELECTRON SHELL) WITH AN APPENDIX TO VOLUME I, PARTS 1, 2, AND 3
Eucken, A.; Hellewege, K.-H. (Editors)
Berlin, New York: Springer-Verlag
1951
724p
In German
This section of the Landolt-Bornstein tables contains data on the electron shells of molecules. The volume includes data published through September 1951. The following properties have been tabulated: band spectra of diatomic molecules, electron band spectra of multiaxial molecules, light absorption of solutions in the infrared and visible region, ionization energies of molecules, optical rotation, molecular rotational, and electronic and optical polarizability of molecules, magnetic moments of molecules, diamagnetic polarizability of molecules, and quantal output of photochemical reactions.
This section contains data on the atomic physical interpretation of crystal physics as of November 1954. The following properties have been tabulated: the characteristics of symmetry, elements, the 32 crystal classes, space groups, lattice types, structures, crystal dimensions of inorganic and organic crystals, atomic and ionic radii, lattice energies, internal vibrations of crystals, electron emission, energy bonds. X-ray spectra and bonding states, electron spectra of crystals, and high frequency spectra of crystals.

This section of the Landolt-Bornstein tables contains tables on atomic nuclei and elementary particles. These tables were compiled by the authors from literature on these subjects published through November 1951. The properties tabulated are: the hyperfine structure of atomic terms and atomic lines, a table of all atomic nuclei and their properties, mass spectrographic measurements, nuclear reaction energies, electron line spectra of artificially radioactive atoms, the radioactive families, nuclear level schemes of radioactive transformations, energy schemes of the light atomic nuclei, beta-spectra, common naturally radioactive atoms and their alpha- and beta-spectra, scattering of charged particles by atomic nuclei, nuclear transition products, passage of alpha-particles, protons and gamma-rays through matter and cosmic radiation.

This section contains 22 mechanical-thermal constants for the equilibrium of heterogeneous systems. Vapor pressure under various conditions, literature references, density, fusion and allotropic changes, temperature changes of crystalline liquids and other data are given for one component systems. The same type of data as well as heterogeneous equilibrium, freezing point lowering, Lewis II: atomic functions, boiling point elevation, osmotic pressure diagrams and references are listed for 222 multi-component systems.

This section of the Landolt-Bornstein tables contains information on solution equilibria of gaseous substances in liquids, solid and liquid metals and alloys, as well as the solubilities of solid and liquid substances in liquid solution media. Except for the section "gases in metals", only solution equilibria in the temperature region of about -200°C to 300°C are presented. The data were taken from literature published through 1962. Solution equilibria are presented for gases with water, aqueous solutions, non-aqueous inorganic liquids, liquids and mixtures of organic and inorganic liquids. Also, solution equilibria of hydrogen, oxygen and nitrogen in pure metals are presented. Solution equilibria are presented for inorganic materials in water and other inorganic liquids, organic liquids, organic materials in water, and solutions with organic solvents.

This section of the Landolt-Bornstein tables includes data on solution equilibria of multi-component systems for solutions of solid and liquid organic materials in organic liquids and equilibria in systems with several non-miscible phases.

This section of the Landolt-Bornstein tables contains data on fusion equilibria and interfacial tensions. The section includes data from literature published through September 1955. Where several values for one measurement were found, the average was presented. Data on fusion equilibria are presented for binary and ternary systems of inorganic compounds, silicate systems, organic systems, and organic-inorganic systems. Interfacial tension data are presented for pure liquids towards their vapors or air, solutions towards air, two non-miscible systems, and interfacial films on water. Data are also presented on absorption from the gas phase and the liquid phase, and on paper chromatography.
#6500532 *0.1151-8.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME II: PROPERTIES OF MATTER IN ITS AGGREGATED STATES, PART 4: CALORIC QUANTITIES OF STATE Schaefer, K. Lax, E. (Editors) Berlin, New York: Springer-Verlag 1961 865p In German

This section of the Landolt-Bornstein tables contains data on the thermodynamic state functions of alloys, compounds and elements, as well as reaction enthalpies for the formation of compounds from the elements and heats of reaction for several important reactions, such as acid-base neutralization and moistening of solid surfaces with liquids. The numerical values are given in joules, except that certain tables for the theoretical calculations of functions present values in joules and calories. Because of the large number of substances and systems listed, numerical data could be given only for a selection of compounds; preference was given to substances which had been accurately investigated or which could claim a greater interest on the part of the user. The following quantities are tabulated: experimental and theoretical basic quantities of thermochemistry, standard value of molar heats, entropies, formation enthalpies and free formation enthalpies; values of thermodynamic properties as a function of temperature for selected materials; heat capacity as a function of temperature; tables for calculating thermodynamic functions in the planes of internal molecular vibrations (including Planck-Einstein and Debye functions); Joule-Thomson effect; magnetocaloric effect of lowest temperatures on paramagnetic salts; thermodynamic data of metal solutions; adsorption heats and heats of moistening; and heats of neutralization.

#690060 *0.1151-5.151-8.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME II: PROPERTIES OF MATTER IN ITS AGGREGATED STATES, PART 5b: TRANSPORT PHENOMENA II, KINETIC HOMOGENEOUS GAS EQUILIBRIA Schaefer, K. (Editor) Berlin, New York: Springer-Verlag 1968 397p In German

Part 5b of Volume II contains some kinetic properties of the elements and selected compounds. The first series of tables lists diffusion constants for elements and ions in metals, salts, and alloys, both liquid and solid. A short table of values for diffusion constants of some gases in selected metals is included. The second series of tables deals with thermal conductivity data for gases, liquids and solids. The constants are reported as a function of temperature, pressure and composition. The next series of tables deals with thermal conductivity data for binary and ternary gas mixtures. Constants are also reported in liquid mixtures and solutions (Ludwig-Soret Effect). The next section lists a series of kinetic rate constants for reaction in gas phase and includes: dissociation, atom-radical and ion-molecule reactions. Another series of tables lists rate constants and equilibrium constants for reactions in solid and mixed phase. Several addenda to the main sections are appended.

This section of the Landolt-Bornstein tables contains data on the electrical properties of matter (except electrochemical systems) published through February 1959. The properties tabulated include the conductivities of pure metals and certain alloys at 0 °C and their temperature variations, resistance ratios, effect of magnetic fields on resistance, super-conducting metals and alloys and their transition temperature, magnetic field effects on super conductivity, galvano-magnetic and thermomagnetic effects, ionic conductivity in crystals, transfer numbers in solids. Also tabulated are the constants of homogeneous semiconductors; photoelectric conduction; piezoelectric equations; and the elastic, piezoelectric and dielectric constants of piezoelectric crystals; Dielectric properties of inorganic and organic crystals; crystalline fluids and pure fluids; dielectric properties at high frequencies; temperature and pressure dependence of dielectric constants; and the dielectric properties of gases. Thermionic emission and work functions, thermopotentials, Peltier and Thomson heats, and photoemission data are also listed.

#6500534 *0.1151-8.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME II: PROPERTIES OF MATTER IN ITS AGGREGATED STATES, PART 7: ELECTRICAL PROPERTIES II (ELECTROCHEMICAL SYSTEMS) Hellwege, K.-H.; Hellwege, A.M. (Editors) Berlin, New York: Springer-Verlag 1960 814p In German

This section of the Landolt-Bornstein tables contains data on the electrical properties of electrochemical systems. The data were taken from the literature published through October 1960. The data compiled include the electrical conductivity of molten salts, pure liquid, and aqueous electrolyte and field solutions; frequency and field strength dependence of conductivity of aqueous solutions; conductivity of ions in aqueous solutions; electrical conductivity of non-aqueous solutions; electrophoretic potentials and electrokinetic potentials, standard electrode potentials, dissociation constants of inorganic and organic compounds and acid-base indicators.

#6500533 *0.1151-3.151-4.151-8.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME II: PROPERTIES OF MATTER IN ITS AGGREGATED STATES, PART 8: OPTICAL CONSTANTS Hellwege, K.-H.; Hellwege, A.M. (Editors) Berlin, New York: Springer-Verlag 1962 901p In German

This section of the Landolt-Bornstein tables contains tables and figures for the optical properties of substances. Data are compiled from literature published through June 1962. Data are compiled for: the optical constants and reflection of metals, alloys, and thin metal layers;
re refractive indices of inorganic and organic materials as a function of wavelengths; optical constants of selected solid materials in broad wavelength regions; rotation of the polarization plane in crystals; double refraction; piezoelectrical and electrooptical constants of piezoelectric crystals; light refraction by glasses and transparent plastic; light permeability; double refraction, gyrotropy; Kerr effect and Kerr effec, refractive index of crystalline fluids; refractive index and optical rotatory power of pure liquids and solutions; Cotton-Mouton effect of liquids; double refraction due to current in liquids; Kerr constants of liquids; refractive indices of gaseous elements and compounds; refraction and dispersion of air and of hydrogen compounds; Cotton-Mouton effect of gases and Kerr constants for gases.


This section of the Landolt-Bornstein tables contains tables and curves of magnetic data for paramagnetic, ferromagnetic, ferrimagnetic and antiferromagnetic substances taken from the literature published through September 1962. The following quantities have been tabulated: fundamental magnetic properties (dia- and paramagnetic susceptibility, spontaneous magnetization, and Curie temperatures) for metallic elements and alloys, exchange energy, crystal energy, magnetostiction, magnetoelectric effects, Faraday effect, magnetic relaxation, Kerr, magnetoelastic effects and resonance in ferromagnetic metals and alloys, properties of ferrospinels, magnetic garnets, perovskites and hexagonal ferrites, magnetic properties of transition elements compounds, lanthanide series compounds and actinide series compounds, paramagnetic centers in crystals, paramagnetic ions as impurities, magnetic susceptibility, cyclotron resonance, de Haas-Schubnikov effect and Faraday effect in semi-conductors, paramagnetic relaxation and nuclear magnetic relaxation.


This section of the Landolt-Bornstein tables gives the paramagnetic and diamagnetic properties of substances. The compiler has listed one or more values for each substance, and marked with an asterisk those considered acceptable and reliable. These, and other selected values considered worthy of inclusion, are listed under "Used References" and values far outside the others under "Further References." The values were taken from literature up to April 1966. Meier magnetic susceptibilities are listed for compounds, and magnetic mass susceptibilities for elements. Diamagnetic susceptibilities are listed for isotropic organic and inorganic substances, and principal susceptibilities for anisotropic substances. Paramagnetic susceptibilities are listed for organic compounds only.

#6500537 10.1151-2.2051-5.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME III: ASTRONOMY AND GEOPHYSICS Bruggencate, J.; Bruggencate, P.T. (Editors) Berlin, New York: Springer-Verlag 1952 795p In German

This volume of the Landolt-Bornstein series contains data on astronomy and geophysics. The material has been taken from literature published through 1951. The astronomical section includes data on astronmetrical instruments, position and time determination, abundance of the elements in the universe, the sun, mechanics and physics of the planets, moons, comets and meteors, stellar radiation, stellar spectra, stellar positions and movements, double stars, variable stars, novae, the galaxy, star clusters and extragalactic nebulae. The section on geophysics contains data on the earth's gravity and tidal forces, radioactivity density, elastic, electrical, magnetic and thermal properties of minerals and rocks; seismicity and earthquake waves; properties of the earth's interior, surface and magnetism; physical and chemical properties of sea water; sea tides, hydrography, meteorology and climatology; and upper atmosphere physics.


This section of the Landolt-Bornstein tables contains material values and mechanical behavior of non-metallic substances, and their technical applications. Data are compiled from literature published through autumn 1964. The following general information is included: measuring systems, units and conversion tables, atomic weights, reduction to standard states, densities of water and mercury. Physical and mechanical properties of non-metallic solids are given for natural and synthetic building materials, organic natural materials (wood, paper, cellulose, wood pulp), fibrous materials, ceramics, glasses, synthetic materials, and natural synthetic materials. Also presented are data on friction and rolling resistance of non-metallic and metallic substances, viscosity, flow through tubes, circuitous flow of bodies during two dimensional flow and rough flow, fluctuation, speed of sound, sound radiation and absorption, the human ear, sound spectra, vibrational numbers, spatial and building acoustics, sound recording and biological effects of ultrasonics.


This section of the Landolt-Bornstein tables includes data on metallographic principles and concepts, testing properties, and the properties of
iron and its alloys. Data are given on diagrams of state, strength testing and destruction free materials. The physical properties of iron tabulated are lattice constants, density, thermal expansion, diffusion, specific heats, enthalpies, specific free enthalpies, phase diagrams and transition points, thermal conductivity, magnetic properties, electrical constants and excitation frequencies. Composition, treatment, strength values and special chemical properties of pure iron, unalloyed steels, highly alloyed steels, special steels, and cast iron are presented.

This section of the Landolt-Börnstein tables deals with the technological aspects of electricity, light and X-rays. It is compiled from the literature published through 1956. Wherever possible the tables have been drawn from several values available for a given quantity, the values they feel is most accurate. The properties tabulated in the electrical section are electrical conductivity, thermal coefficient of resistance; conductivity of molten metals; effect of pressure, tension, and cold machining on electrical resistance; resistance as function of frequency; resistance of thin layer semiconductors; thermoelectric emf of metal pairs and thermocouples in use; effects of pressure and cold treatment on thermocouples; conduction of electricity in gases, breakdown potentials of gases; insulating liquids, filling and impregnating media, fibrous material insulators, insulating foils, mica insulators, rubber insulators; resin, plastics, quartz, glass, ceramic materials and rocks as insulators; and magnetic materials. The quantities tabulated in the light section are: sensitivity of the human eye, illuminating materials, light sources, light filters and photographic films. The X-ray technology includes data on measurements and units of X-ray technology.

This section of Landolt-Börnstein is a compilation of data dealing with the technology of thermal measurements and the thermodynamic properties of gases, vapors, liquids, and solid materials. Data are included for such areas as the basis of temperature measurement; the temperature measurement with contact thermometers; optical temperature measurement (Pyrometry); x-ray diffraction; and thermodynamic properties of homogeneous materials, gases, water and steam, refrigerating agents; thermal expansion of solids and liquids; and the specific and thermal properties of technically important materials.

This is the third part in a series of Landolt-Börnstein tables presenting the physical properties and behavior characteristics of most of the elements in the periodic system exhibiting metallic characteristics as well as data concerning their alloys. Special emphasis is directed toward light metals, special work metals, binding materials; enamel properties, corrosive behaviors, reactor metals, metals of the rare earth, semiconductors, and current developments in binary alloy systems. The volume is divided into sections entitled: titanium, beryllium, aluminum, magnesium, lithium, rubidium, cesium, liquid metals; nuclear survey of reactor metals, uranium, plutonium, zirconium, hafnium, thorium, rare earths, semiconductors, bearing metals, thorium, solder, welding and cutting metals, metal adhesives, and corrosive behavior of materials.

This first volume of the New Series of Landolt-Börnstein contains excitation levels and decay schemes for all nuclei from A = 5 to A = 257. The essential parts of the material are the level and decay schemes. These are supplemented by tables and special data. The tables are the following parameters for each level: excitation energy above ground state; the total angular momentum quantum number and the parity; the isobaric spin quantum number; the half life or mean life of the state, its half-width; the known modes of decay of the states; binding energies for various nucleon combinations, and the Q-values for the relevant nuclear reactions. Much of the same information is incorporated and in diagrams in which the energy levels of the various nuclei are exhibited schematically, together with the reactions by which they are formed. Recent
The volume contains nuclear radii and other characteristic parameters of atomic nuclei which are deduced from electron scattering experiments and the observation of mesic X-rays. The first section contains data on nuclear charge distributions derived from sources other than scattering, primarily mu-mesic atoms for spherical nuclei. Data are given on energy levels and radius parameters for mu-mesic X-ray transitions; also for neighboring nuclides, deformed nuclei, and radii of isomers. The second section deals with nuclear radii determined by electron scattering. Both electric charge and magnetic radii are given. Because a variety of nuclear models are available, each nucleus is discussed separately, in addition to listing the size parameters.

The various types of angular correlation coefficients have been tabulated. These tables provide the coefficients for a complete calculation of the basic angular correlations in alpha-, beta-, and gamma-spectroscopy. In chapter 2, the angular correlation formalism is given in a necessarily incomplete collection of relations involving the tabulated coefficients, i.e., 3j-symbols, 6j-symbols, F-coefficients, and GAMMA-coefficients. In the computer tables these symbols are given in upright capital letters: J = j, J' = j' etc., and GAMMA = Y. These quantities are given solely by angular momentum conservation for the total system considered. When answering the question of how the angular momenta of the various subsystems of particles may combine to fulfill the conservation condition, one encounters with Clebsch-Gordan-coefficients for two angular momenta and with Racah-coefficients for more than two. To achieve a higher degree of symmetry, Wigner has rewritten both types of coefficients as the 3j- and 6j-symbols, respectively. In more complicated correlations (e.g., triple correlations, where the intermediate transition also is observed) a new combination of 6j-symbols appears which is usually called the 9j-symbol. For alpha-, beta-, and gamma-spectroscopy a combination of one 3j- and one 6j-symbol with statistical factors is frequently used and is called the F-coefficient. The GAMMA-coefficient is a slightly modified form of this F-coefficient useful for beta-spectroscopy. The coefficients mentioned above are extensively tabulated in this volume. For the 9j-symbol, however, only selected ones are listed. In addition, there is a chapter dealing with angle functions. Since angular correlations are generally developed in a series of Legendre polynomials, these functions should be very useful.

These beta-decay tables present certain combinations of electron radial wave function and phase shifts for both screened and unscreened beta-decay functions. The tables for electron capture give the amplitudes and values of the electron radial wave functions at the nuclear radius for the k, l and m shell for all values of l (the l and m shells are excluded for very light nuclei). A brief compilation of all the relevant formulae used in the treatment of beta-decay and electron capture is given. An explanation of the details of the compilations and a description of the tables is included; in addition, 56 references are given.

This book contains data on the magnetic properties of free radicals, including inorganic and organic uncharged free radicals and radical ions. These data were taken from published results of ESR and atomic beam resonance experiments. The data compiled include: name and structural formula of a free radical compound, method of generation, crystal matrix or solvent used for the radical, temperature of measurement, frequency of measurement, g-factor, hyperfine splitting parameter, primary references for the values quoted, and further references on the same subject.
This volume lists data on acoustics based essentially on the molecular properties of matter, and values of sound velocity, absorption coefficients, and dispersion data. The measurements were made at frequencies ranging from 10 kc to 200 kc. Preference was generally given in compiling data to measurements for which statements on density, chemical purity, and the temperature coefficient were added. Often, however, several statements can be found for a measured value. Generally, the limit of accuracy is 1½ to 2%. Data are given on sound velocity in gases and liquids; for pure gases and liquids as a function of temperature, pressure, and structure; also for mixtures of gases, mixtures of liquids, and solutions. Data are compilation on sound absorption and dispersion in gases; sound absorption in homogeneous liquids, in solution, and on sound propagation in isotropic or quasi-isotropic solids (metals, inorganic non-metals, and organic solids). There are special sections on sound propagation in liquid helium and shock wave velocity.

In German and English

This volume of the new Landolt-Bornstein series gives the elastic, piezoelectric, electrooptic, and electromechanical constants of crystals. The data included were taken from data published in the literature through 1965. For materials where one or two sets of measurements were given, the mean and coefficient of variation are given. For non-piezoelectric constants, the elastic stiffness and compliance constants, together with their temperature and pressure variations, are given. For piezoelectric constants, the elastic, piezoelectric, dielectric, piezo-optic, and electrooptic constants, plus the electromechanical coupling factors are given. The temperature coefficients of the elastic stiffness, compliance, piezoelectric stress and strain constants are also given.

In German and English

Volume III/2 of the New Series of Landolt-Bornstein continues Volume III/1 only three years after publication of the latter. All the tables contained in III/1 are brought up to date by supplements in III/2. In addition, the elastic and electrooptic constants of higher order and the nonlinear optical susceptibilities have now been included. Readers should note that the Index of Substances at the end of III/2 refers to both volumes.

In German and English
This volume of the Landolt-Bornstein New Series contains data useful in the fields of astronomy and astrophysics. The following topics are covered: optical instruments, radio-astronomical devices, performance of telescopes, photoelectric photometry, position and time determination, astronomical constants, abundances of the elements in the universe; physical and mechanical properties of the sun, planets, satellites, comets, meteors, artificial earth satellites, interplanetary space, stellar positions and motions, stellar spectra, star colors and magnitudes, physics of stellar atmospheres, stars and evolution, double stars, variable stars, novae and supernovae, planetary nebulae, star clusters and association, properties of the nearest stars, galactic structure, kinematics and dynamic properties of the galaxy, interstellar space, galaxies, and clusters of galaxies.

This handbook consists of mathematical tables and methods, physical and chemical data, reaction kinetics, reactor design, thermodynamics, fluid and particle mechanics, transport and storage of fluids and solids, size reduction and enlargement, heat generation, transport and storage, heat transmission, heat-transfer equipment, refrigeration, distillation, gas absorption and solvent extraction, humidification and drying, absorption and ion exchange, other diffusion operations, multiphase contacting and separations, process control, materials of construction, mechanical plant and project engineering, and cost of profitability estimation.

This book presents a critical review of the various estimation procedures for a limited number of properties of gases and liquids. These include critical properties, P-V-T and thermodynamic properties, vapor pressures, latent heats, heats of formation, free energies of formation, heat capacities, surface tensions, viscosities, thermal conductivities, diffusion coefficients, and equilibrium properties of mixtures. Comparisons of experimental and estimated values are shown in the form of tables for indication of the degree of reliability of the methods discussed. The book is designed to be used as a text for molecular physics or a reference source for engineers. A set of Appendices summarizes material referred to in the chapters and contains a large volume of tabulated data as well as references to the literature.

This compendium was compiled from literature data available to the authors (30 Russian experts). An attempt was made to present the most reliable numerical values; the criteria for reliability were the experimental techniques used to obtain these values, the purity and state of the samples used, and statistical criteria. One recommended
numerical value is given for each property, but additional literature references are listed. The properties included in the handbook are: 1) general information on properties of the elements and atomic structure; relative abundances, atomic weights and atomic volumes, atomic and ionic radii, density, crystal structure, optical and X-ray spectra; 2) nuclear properties of the elements; isotopes, nuclear energy levels, neutron interaction cross sections, and scattering of electrons; 3) thermodynamic and heat properties; entropy, heat capacity, melting and boiling points, heats of fusion and sublimation, vapor pressures, critical parameters, thermal conductivity, crystal lattice energy, surface tensions, viscosity, dielectric parameters, ion mobilities; 4) electrical and magnetic properties; electrical conductivity, superconductivity, thermolectric properties, magnetic properties, emission properties, dielectric properties; 5) optical properties; color, emissive, reflective, refractive index; 6) mechanical properties; elastic and shear moduli, Poisson ratio, compressibility, velocity of sound, tensile, yield and compressive strength, fatigue limit, creep, and hardness; 7) electrochemical properties; electrode potentials, oxidation-reduction potentials; 8) chemical properties; interaction of the elements with various media and other elements.

#6809006  *0.2173
Scher, J. A. (Editor)
HANDBOOK OF BIOCHEMISTRY, SELECTED DATA FOR MOLECULAR BIOLOGY
Cleveland, Ohio: The Chemical Rubber Company
1968
900p

This handbook gives evaluated data for selected areas of biochemistry including the principal research frontiers in medicine, modern biology, genetics, biophysics, and molecular biology. The material is divided into the major sections of abbreviations and nomenclature, amino acids, peptides, and proteins, carbohydrates, lipids, steroids, purines, pyrimidines, nucleotides and oligonucleotides, nucleic acids, genetics and biology, physical and chemical data, and a miscellaneous section.

#6810020  *0.2173
Stecher, P. G. (Editor)
THE MERCK INDEX, AN ENCYCLOPEDIA OF CHEMICALS AND DRUGS EIGHTH EDITION
Rahway, New Jersey: Merck & Co., Inc.
1968
1214p

This eighth edition of the Merck Index includes nearly 10,000 descriptions of individual substances, more than 4,500 structural formulas, and about 42,000 names of chemicals and drugs. References to the cited original papers are given. The Merck Index has a strong medicinal element, but it is essentially an organic chemical work with the inclusion of useful inorganic substances and a separate section on organic name reactions. The Cross Index of Names and Formula Index provide a key to the monographs included. The handbook includes some data and conversion factors in tabular form, however, a majority of the information is given in paragraph form under each alphabetical entry.

#6512008  *0.2172-3.172
TABLES OF COULOMB WAVE FUNCTIONS, VOLUME I
NBS Applied Mathematics Series No. 17, National Bureau of Standards, Washington, D. C.
April 1952
141p
Available from Superintendent of Documents, GPO

This report presents numerical values of the Coulomb wave functions in somewhat peculiar notation. However, an introduction by M. Abramowitz defines all quantities and it is possible to reconstruct the actual Coulomb wave functions from the tables presented herein. In addition, there is a discussion of the significance of these wave functions in a preface by G. Breit.

#6708016  *0.2173
TABLES RELATING TO MATHEW FUNCTIONS
August 1, 1967
311p
Available from Superintendent of Documents, GPO

This second edition of "Tables Relating to Mathieu Functions" (Columbia University Press, New York, 1951) incorporates the article "Table of Characteristic Values of Mathieu's Equation for Large Values of the Parameter" by Gertrude Blanch and C. D. Jerri, Journal of the Washington Academy of Sciences, 45(6), 166-170, June 1955, but makes no essential changes in tables or introductory material. However, a number of misprints in the first edition have been corrected and the bibliography updated.

#6500541-6500548, inclusive  *0.1083
Washburn, E. W. (Editor)
INTERNATIONAL CRITICAL TABLES OF NUMERICAL DATA, CHEMISTRY AND TECHNOLOGY VOLUMES 1 TO 7 AND INDEX
1926-1933
1-415p; 2- 616p; 3- 444p; 4- 481p; 5-465p; 6- 471p; 7- 507p; Index- 321p

The International Critical Tables, covering the literature from 1910-1923, provide evaluated data on physical, chemical, and technological materials. The seven volumes of data are organized into more than 300 sections. Among the sections and properties included are national and local systems of weights and measures, symbols, basic constants, conversion data, instrumentation data, chemical elements and atoms, thermometry laboratory techniques, laboratory methods for producing and maintaining constant temperatures, constant humidity, pressure, volume of liquid mercury, weights and weighing, calibration of volumetric vessels, buffer solutions and indicators, high vacuum technique, building material properties, abrasives, glass, fuels and petroleum, paints and varnishes, X-ray diffraction data for industrial materials, metallurgy, liquid and gas kinetics, spectroscopy, photometry, photography, properties of soaps, electrical conductivity and resistivity, magnetism, acoustics, refractivity, and optical properties, explosives, PVT relations for one-phase systems, phase equilibrium data, osmotic pressures, surface properties, and a great volume of other related areas and properties. The last volume is a comprehensive index of all materials and properties listed in the International Critical Tables.

#6711003  *0.2183
Weast, R. C. (Editor)
HANDBOOK OF CHEMISTRY AND PHYSICS, 49TH EDITION
Cleveland, Ohio: Chemical Rubber Company
1968
2450p

This Handbook is a compendium of data from all fields of physical science which have been experimentally determined and published by scientists over the years. The Handbook is a collection of tables of important mathematical functions. The mathematical tables include logarithmic and trigonometric functions, numerical tables of squares, cubes, powers, roots and reciprocals, statistical tables, derivatives and integrals, algebraic functions, and other formulas. Data on elements and inorganic compounds includes atomic weights, electron configurations, neutron cross sections, and physical constants. Data on organic compounds include naming rules,
physical constants, organometallic physical constants, properties of steroid hormones, amino acids, carbohydrates, and waxes. General chemical data includes thermodynamic constants, dissociation and ionization constants, conductances, electrolytes, melting and boiling points, vapor pressure, and properties of aqueous solutions. General physical constants data includes thermal conductivity, dielectric properties, superconductivity, magnetic properties of matter, X-ray and optical spectral index of refraction.

#6809004  *0.2183
Wick, O. J. (Editor)
PLUTONIUM HANDBOOK, A GUIDE TO THE TECHNOLOGY, VOLUME 1

The Plutonium Handbook provides information on the range of topics concerning plutonium technology. The book is divided into seven separate sections: physics; metallurgy and ceramics; chemistry; chemical separations; fabrication and utilization; analysis and inspection; and health and safety. References are included and data are given both graphically and in tabular form.

#6603015  *0.2183
Wilhoit, R. C.; Hathaway, W.
TABLES OF CONVERSION FACTORS BASED ON ACCEPTED CONSTANTS AS OF 1965
College Station, Texas: Texas Agricultural and Mechanical University, Chemical Thermodynamic Properties Research Center, College Station, Texas November 1965 70p

These tables contain values of the defined, basic and derived physical constants, and tables of conversion factors for physical quantities measured in different units, presented in matrix form.

#6711019  *2.683-3.183-0.2183
Wilson, B. S. (Editor)
THE RADIOCHEMICAL MANUAL, SECOND EDITION
Amersham, England: Radiochemical Centre 1966 327p

The manual is intended as a guide for all who use radioactive substances professionally; its aim is to help the individual user choose the right material for his purposes and to make effective use of it. The text portion of the book includes information on preparation of radioisotopes; radiation sources; problems of using radioactive materials; radioactive chemicals as pharmaceuticals; quantification of radioactive substances; radiological safety and waste disposal; packaging and carrying radioactive materials; ownership and use regulations and a guide to literature. The tabulated data includes: physical data on the more common radioisotopes, including physical characteristics, availability, beta and gamma decay energies and half-lives; radiation sources, synthetic routes to labelled compounds, decomposition rates of labelled compounds, measurement and safety.
Nuclear Properties

(Including Fundamental Particles Properties)
The decimal variations (d) of relative atomic weights (m) in the form d = M - A + 1, are graphically represented as a function of the mass number A by 5 linear segments.

The results of an analysis which utilized the experimental differential cross section for elastic scattering of neutrons are presented. The calculated quantities for all materials were the elastic scattering cross section, the average logarithmic energy decrement per collision, and the Legendre expansion coefficients. The neutron energy range was approximately 0.1 to 10 MeV. These parameters were presented graphically as a function of neutron energy when sufficient data were available. For materials in which this was not the case, the results are presented in tabular form.

This initial collection contains benchmark problems relating to numerical determination of space, time, angle, or energy distribution of particles in an assembly. The primary objective is to provide reliable solutions to diverse problems of varying complexity. This book should serve as a source book of solutions to mathematically well-defined problems for which either analytical or very accurate approximate solutions are known.

Two tables are presented, one for protons and one for electrons, which enable the computation of the rate of energy loss, given the value of the mean excitation potential. While the value of this latter constant has been taken, in this paper, as being given by \[ I = a Z = 11.5 Z \text{ eV}, \] the validity of such a simplistic formulation has been increasingly questioned, several recent experiments yielding results considerably different from those expected. The range-energy curves therefore are tentative, pending a better understanding of the mean excitation potential. The energy-loss considered in this paper derives only from collision effects. The losses due to bremsstrahlung and meson and pair production result from an essentially different process. The calculation of the Fermi effect is ill-advised pending a better determination of the constants.
He-3, or He-4 as either incident or product particle for about 650 target nuclides. Approximately 8000 Q values are tabulated.

He-4 stable survey

C Nuclear neutron energies, elastically

Included thermal-neutron-

PHOTONS the product water, an

are mass-spectroscopic

THERMAL-NEUTRON-CAPTURE

65%257-274

Tschalaer, Attention also given

A3:343-360

January

REVIEW

Barashenkov, #6509013

January 1969

This review of the properties of leptons, mesons, and baryons is an updating of Rosenfeld et al. [Rev. Mod. Phys. 40, 77 (1968)]. Data are evaluated, listed, averaged, and summarized in tables and wallet sheets. A data booklet is also available.

Bartholomew, G. A.; Higgs, L. A. 


146p

This is a compendium of energies, absolute intensities, and spectral distributions, together with a complete bibliography, of neutron capture gamma-rays compiled from information available as of June 1, 1958.

Bartholomew, G. A.; et al.


A compilation is presented of thermal-neutron-capture gamma-ray energies and intensities with decay schemes, examples of original data, and fully corrected spectral distributions. A table of stable isotopes with abundances, thermal-neutron radiative-capture cross sections, spins, parities, and other information of value to neutron-capture gamma-ray spectroscopy is included.

Beckurts, K. H.; Wirtz, K. 

Dresner, L. (Translator) 

NEUTRON PHYSICS Berlin and New York: Springer-Verlag 1964

454p

In German

This text presents a survey of: the physical properties of the neutron, neutron sources and detectors, and cross-section measurements; the theory of neutron fields; the theory of diffusion, slowing down, and thermalization; the determination of neutron fluxes and distributions; and the determination of neutron transport properties.

Berger, M. J.

ENERGY DEPOSITION IN WATER BY PHOTONS FROM POINT ISOTROPIC SOURCES Journal of Nuclear Medicine, Supplement 1, No. 2:17-25 February 1968

Buildup factors and related data which describe the deposition of energy by photons in water medium are presented. Nineteen photon source energies between 15 keV and 3 MeV are included. Data are given on energy-absorption buildup factor for point isotropic sources in water, coefficients of polynomial representation of energy absorption buildup factors for water, specific absorbed fractions and absorbed dose rates in water and muscle, and radii of spheres for specific absorbed fractions.

Bertin, M. C.; et al. 

NEUTRON CROSS SECTIONS OF U-238, U-235, U-239, U-234, U-236, Pu-239, Pu-240, W, Pb, Ni, Cr, C, Li-6, Li-7, AND T

UNC-5099, United Nuclear Corporation, Division White Plains, New York December 31, 1964

303p

Available from CFSTI, AD 616629

Neutron cross-section sets have been prepared for 16 elements or isotopes for neutron energies from 0.037 eV to 18 MeV. The cross sections tabulated include the total, elastic, inelastic, (n,2n), and fission cross sections, as well as cross sections for charged particle emission. Information is also given on the angular distribution of elastically scattered neutrons and on the energy distribution of neutrons and gamma-rays following nonelastic scattering.

Bhanot, V. B.; Gupta, S. C. 

MASS-SPECTROSCOPIC ATOMIC MASS DIFFERENCES Nuovo Cimento, 10:1011-1023 June 1963

New mass-spectroscopic data on atomic mass differences in the medium and heavy mass regions were compiled and tabulated. Corresponding values in the new scale based on C-12 as the standard are also given. Unadjusted experimental values for nuclides in the new scale were listed. For a large number of stable nuclides in the region from strontium to antimony the newer values obtained from the data of Demirkanov et al., are more accurate than the corresponding values listed in the recent exhaustive mass table of Koenig et al. Attention is drawn to the advantages of the isotopic doublets over the published conventional mass doublets in the heavy mass region.

Bichsel, H.; Tschaaler, C. 

A RANGE-ENERGY TABLE FOR HEAVY PARTICLES IN SILICON Nuclear Data, A3:343-360 October 1967

Stopping-power S and range R tables for protons, deuterons, tritons, He-3, He-4, and Li-7 ions of energies between 1- and 200-MeV projecting silicon absorbers are presented. Auxiliary data are included in tables for the coefficients C and alpha needed for the approximate expression S=0.6456 and in figures giving the energy dependence for the coefficients of R = C x E^alpha. Included is a short review of principles to be considered in detector applications.

Bormann, M. 

NEUTRON SHELL EFFECTS IN THE (n,2n) CROSS-SECTIONS AT 14 MeV Nuclear Physics, 65:257-274 May 1965

The systematics of the available total (n,2n) reaction cross sections for 14 MeV neutrons were investigated for even-neutron target nuclei. The existence of neutron shell effects is verified and discussed on the basis of the statistical theory.
Minima in the \((n,2n)\) cross sections at the magic neutron numbers \(N = 28\) and \(50\) for even- and \(N = 20\) for odd-proton nuclei can be accounted for by similar trends in the \(Q\) values. Cross-section maxima at the closed neutron shells \(N = 82\) and 126 for even- and \(N = 28, 50, 82,\) and 126 for odd-proton target nuclei can be understood as an effect of shell closure on the nuclear level density, the relative decrease of which on passing across a closed shell is larger for higher excitation energies. The \((n,n'\gamma)\) reaction is the principle one competing with the \((n,2n)\) process at 14 MeV in the medium and heavy mass regions. Since for the \((n,n'\gamma)\) reaction the residual excitation energy lies in the region up to 12 MeV, whereas for the \((n,2n)\) reaction it does not exceed 4 MeV, and since the relative change in the level density is greater at the higher excitation energies, the competition between \((n,n'\gamma)\) and \((n,2n)\) reactions favors the latter in the region of closed neutron shells and thus accounts for the observed maxima in the \((n,2n)\) cross-section systematics. Ninety-six references are included.

---

Available from AEC Dep. Libraries

The steady-state distributions of electron energies in helium, neon, argon, krypton, and xenon are given at values of \(E/N\) (ratio of dc electric field to gas atom number density) up to \(E/N\) approximately equal to 10-17 V/sq.cm.

---

Cross sections based on new values (given) for the cross sections of the monitor reactions \(Al-27(p,3pn)Na-24\) and \(C-12(p,n,p)nC-11\) are tabulated for the production of specific nuclei in proton reactions on nuclei from Be through U. Cross sections for production by gamma reactions on \(B\) and \(Ni;\) deuteron reactions on \(Cu,\) \(Th,\) \(U,\) \(Ar,\) \(Ne,\) \(Mg,\) \(N,\) \(C,\) \(and\) \(Al;\) alpha reactions on \(U,\) \(Th,\) \(Al,\) and \(C;\) \(C-12\) ions on \(Au\) and \(Al;\) and by \(Li-6, N-14,\) and 0-16 ions on \(Al\) are also included.

---

The tables list kinematic quantities for two-body and quasi two-body photoproduction reactions. Although the laboratory angle and momentum are given for both outgoing particles, differential quantities such as the solid angle transformation, Jacobians, etc., are given only for the long-lived charged particles (\(N, K^*, p,\)) to this extent the tables are oriented toward spectrometer experiments. Each table is characterized by the laboratory photon energy shown in the upper right-hand corner. Tables are not given for photon energies below the kinematic threshold for the reaction. Each table lists the various kinematic quantities every \(5^\circ\) in the center of mass from \(0^\circ\) to \(180^\circ.\)

---

A compilation is presented of formation cross sections in the region beryllium to copper for nuclear reactions above 50 MeV.
A Fortran code has been developed for the calculation of eigenvalues and eigenfunctions for neutrons in real potential wells. A systematic procedure is given for approximate location of the eigenvalue and an automatic search procedure to determine the exact location. The code may be used for either bound or scattering states. In the case of scattering states, the criterion for maximum scattering (80° phase shift) is used to determine the energy of the state. The eigenvalues are determined by matching the numerically calculated logarithmic derivative (f1) inside the nucleus to the appropriate analytical logarithmic derivative for the region outside the nucleus. In an alternate mode of operation, the outside value of f1 may be set arbitrarily, and a match made to this value. Sample results for a Woods-Saxon well with spin-orbit coupling for the case of oxygen-16 are shown. The code is in Fortran and was written for an IBM 7090 computer.


This revised edition of the Handbook of Criticality Data, AHSB(S) Handbook I, contains the best estimates on the critical parameters of the most reactive systems likely to be encountered in process plants. It presents data on compounds of plutonium and uranium, and some account is taken of the thermal poisoning effect of the nitrate and of Pu-239. Metal-water data and unreflected data are included in this edition. No theoretical justification of the data is given, nor is correlation with experiment shown on those graphs where the information is firmly established. In cases where sufficient experimental evidence or data check points are not available, graphs are overprinted as provisional. The basis of calculation is given and quantitative comparison with other known calculations is quoted as appropriate.


A comprehensive compilation of the data available through December 1963, on total cross sections for the (n, alpha) reaction on nuclei (64 < A < 238) at 14 MeV is presented.

CINDA 68 AN INDEX TO THE LITERATURE ON MICROSCOPIC NEUTRON DATA CINDA 68, Atomic Energy Commission, Division of Technical Information Extension, Oak Ridge, Tennessee 1968 1017p Available from AEC Div. of Tech. Information, TID 24480

CINDA 68 (Computer Index of Neutron Data) is a cumulative bibliography of the literature on microscopic neutron cross sections and allied data for any given target nucleus, arranged first by element and mass number, then by cross section or other quantity. It is intended to indicate where in the journal and reports literature information can be found on specific microscopic neutron data and is not a listing of the data themselves. CINDA is limited to reactions induced by neutrons of energy less than 20 MeV for specific elements and isotopes and includes information on (gamma,n) and (gamma,f) reactions with the restriction that the gamma-ray energy must be less than approximately 15 MeV and the (gamma,n) cross section greater than 0.1 mb. CINDA 68 contains the full computer file as of June 15, 1968.

CINDA 69 AN INDEX TO THE LITERATURE ON MICROSCOPIC NEUTRON DATA CINDA 69, Atomic Energy Commission, Division of Technical Information Extension, Oak Ridge, Tennessee May 1969 1157p Available from AEC Div. of Tech. Information

The CINDA (Computer Index of Neutron Data) bibliography allows its users to find the references to specific types of cross section information or other microscopic data from neutron-induced reactions, for any given target nucleus. CINDA entries are ordered in this publication first by element and mass number, then by cross section or other quantity. Within these isotopes and quantity groups, the references are ordered by date of publication. Editors should note that experimental, theoretical and evaluation articles are mixed within the listing.

CINDU CATALOGUE OF NUMERICAL NEUTRON DATA AVAILABLE FROM THE IAEA NUCLEAR DATA UNIT CINDU-6, International Atomic Energy Agency, Nuclear Data Unit, Vienna, Austria November 1967 116p Available from the IAEA Nuclear Data Unit

This issue of CINDU (Catalog of the IAEA Nuclear Data Unit) contains a bibliography of the current holdings of the data unit as of 1 November 1967. Only those references stored in the data storage and retrieval system (DASTAR) are listed in CINDU. (CINDU primarily lists papers on neutron induced and neutron scattering reactions). For each reference, there is listed the name, atomic and mass number of the target nucleus, the quantity measured or calculated, the energy range considered, the publishing laboratory, the type of article (experimental, theoretical, etc.), journal reference, volume, page and date, the author's name, and an accession number to the DASTAR data from the IAEA Nuclear Unit.
The rays emitted by radioisotopes produced in the thermal neutron activation of natural isotopes are tabulated according to energy. The natural abundance of the isotope, its half life, thermal neutron cross section, and absolute intensity of the radiation are all given. The source of each energy tabulation is given. Ninety-one references are included.

**Photon energies and abundances were compiled and summarized for some fission-product and other radio-nuclides, using data reported in the literature up to June 1963. The data are presented in tabular form, listing photon energies and abundances for gamma rays, beta rays, and X-rays emitted. A list of multipliers is also presented for converting activities of the radioisotopes to infinite-plane exposure rates.**

**Atomic mean excitation energies (I values) have been obtained from stopping-power and range data in twelve experiments. The analysis incorporates new values of shell corrections that include contributions made by electrons in all shells. Except for one set of data, the values of I obtained from different experiments are consistent with one another. Empirically, it is found that the following formula fits the I values obtained here:**

\[
I(\text{eV}) = 11.2 + 1.17 Z, \quad 2 \leq Z < 13, \\
I(\text{eV}) = 52.8 + 8.71 Z, \quad 2 > Z. 
\]

The numerical values found from the experiments analyzed are in agreement with those suggested in the National Academy of Sciences-National Research Council Report 1133. It appears that a few of the I values given in the National Bureau of Standards Handbook 79 (1961) are now out of date. A table of I values for some thirty-six materials of particular importance in radiation dosimetry is given in addition to the elemental I values found from the new data analysis.

**All measured Fermi matrix elements of normal allowed beta decay are compiled. For each case an adopted value is given, which is the mean overall consistent results. The Fermi matrix elements turn out to be small or vanishing, in fair agreement with the isospin selection rule. The question of charge dependence of the nuclear forces is discussed.**

**This volume of the proceedings includes all of the invited papers, formal comments and discussion of the papers presented in the third Symposium on the structure of low-mass nuclei held at the University of Kansas on April 18-20, 1968. The papers discuss such topics as the studies in the 2s-1d shell using triton induced reactions; effective interactions in nuclei and two-nucleon scattering; nuclear forces and the structure of light nuclei; gamma decay of analogue states in the 2s-1d shell; T = 2 levels in Ne-20, Mg-24, and S-32 as compound nucleus resonances; the mirror nuclei, Mg-25 and Al-25; gamma decay of states in Al-25; shell-model calculations in the 2s-1d shell; theory and experiment in the 2s-1d shell nuclei; SU3 symmetry and realistic interactions; gamma-ray spectroscopy in light nuclei; Na-22; recent developments on the dynamitron accelerator; and heavy ion acceleration.**
prepare a revised data file DFN 50 for inclusion in the U.K.A.E.A. nuclear data library.

#6604066  *2.085-2.2783*  Dragoun, O.; Pauli, H. C.; Schmutzler, F.  TABLES OF INTERNAL CONVERSION COEFFICIENTS FOR N-SUBSHELL ELECTRONS  Nuclear Data Tables, A6:235-351  September 1969

Internal conversion coefficients are presented for the first five N-subshells for every value of the atomic number from 2 = 60 to 2 = 100. The coefficients are given for nuclear transition multipoles E1,...,E4, M1,...,M4 and for transition energies from about 1 keV above threshold to usually 500 keV. The finite nuclear size is taken into account as well as atomic screening according to the nonrelativistic, self-consistent field method.

#6508047  *2.323.*  Dunner, J. E., Jr.  LOS ALAMOS HANDBOOK OF RADIATION MONITORING, THIRD EDITION  L.A.-1835  Los Alamos Scientific Laboratory, Los Alamos, New Mexico  November 1958  180p

Available from the Superintendent of Documents, GPO

This third edition of the Handbook of Radiation Monitoring contains data for permissible exposures and permissible body burdens, as well as numerous tables, graphs, and tabulated information on the basic data needed frequently in a nuclear laboratory. This manual contains a monitor's check list, radio code, emergency monitoring, protective methods, LASL radiation safety policies, permissible dose and measurement units, permissible contamination levels, film examination, table of permissible exposures, table of isotopes, monitoring instruments, alpha monitoring, beta-gamma monitoring, neutron monitoring, tritium monitoring, personal dosimetry, urinalysis, air sampling, and respiratory protective equipment information.


This is a preliminary report to the National Research Council consisting of a reevaluation of the atomic constants and conversion factors of physics. It is of historical interest only. As a result of the precision microwave and atomic beam techniques, it reported advances in the accuracy of such constants as e and m, the charge and mass of the electron, h, Planck's constant, Avogadro's number, the velocity of light, the conversion factor from the Siegbahn nominal scale of x-units to milligrams, and a list of some sixty or more useful derived constants and conversion factors which can be computed from the preceding with the use of accurately known other constants, such as the Rydberg, the gas constant, etc. In this report the least-squares method was employed to adjust the data to arrive at a set of consistent values for the constants in question.

#6509034  *2.1623*  Dutta, A. K.; et al.  ON NUCLEAR BINDING ENERGIES  Indian Journal of Physics, 38:57-60  January 10, 1964

Relations for the binding energies of nuclei of a given mass number were derived. Experimental and calculated binding energies for all the isobars of some mass numbers are tabulated.


Available from CFSTI

Gamma ray production cross sections are presented as a function of 15 neutron lethargy groups and 11 photon energy groups for 15 elements and isotopes. These data were compiled by Proctor GAMMA-P and account for the production of gamma rays by neutron radiative capture, neutron inelastic scattering, neutron induced fissioning, and (n, alpha) reactions.

#6907025  *2.2552*  ENDF NEWSLETTER 3:MAT1001-1 to MAT1072-3, Brookhaven National Laboratory, Upton, New York  October 1967  100p

This newsletter contains a summary of neutron cross section data, information and references for H, D, Li, Be, B, C, N, O, F, Ne, Mg, Ti, V, Mn-55, Mo, Xe-135, Sn-114, Cu-64, Zn-64, Cd, In-115, Lu-178, Lu-176, Au-197, U-235, Pu-238, Pu-239, Pu-240, Pu-242, Cm-244, Na-23, Hf-176, Hf-177, Hf-178, Hf-179, Hf-180, Hf-174.

#6500044  *2.1423*  Endt, P. M.; Van der Leun, C.  ENERGY LEVELS OF LIGHT NUCLEI III.  Z = 11 to 2 = 20  Nuclear Physics, 341-324  1962

The present compilation of information is for the energy levels of nuclei with Z = 11 to Z = 20. The nuclei are presented in order of increasing Z, and nuclei of the same Z, in order of increasing A. Generally, each nuclear reaction is treated under the heading of the final nucleus. Exceptions are reactions where resonances have been observed; and the beta decay of unstable nuclei. In all other cases, the order of the reactions is determined by the initial nucleus, starting with the lowest Z element. Following the last reaction discussed, for each nucleus a list is given of reactions "Not Reported". Discussions on the energy levels of a particular nucleus with depictions under the headings of a specific reaction leading to that nucleus have been given as "Remarks" at the end of the survey of that nucleus. References are given for all the information provided in the text, tables or figures. The excitation and resonance energies given in the level diagrams are the weighted mean values of all determinations available. The near maximum excesses used throughout this compilation are listed in table 2. The natural abundances of the isotopes in the mass range discussed in this compilation are listed in table 2, together with nuclear moments.

#6807002  *2.1423*  Endt, P. M.; Van der Leun, C.  ENERGY LEVELS OF Z = 11-21 NUCLEI (IV)  Nuclear Physics, A105:1-488  December 1967

A compilation is given of experimentally determined properties of energy levels of Z = 11 to 21 nuclei with special emphasis on nuclear spectroscopy. Data are presented such as excitation energies, half-lives, and (n, gamma) capture cross sections. All available elementary particle resonances are reported and updated values given. Over 3200 references are provided.
The experimental data on alpha energies, half-lives and branching ratios are presented in tables. An attempt has been made to make the calibration energies of various measurements compatible. The experimental alpha-decay energies and those given by the 1964 atomic mass table and mass formulae are presented graphically as a function of the neutron number of the parent nucleus.

These Q-values have been computed from masses of nuclides participating in the reaction, using a consistent set of nucleonic masses derived from all the relevant experimental data, this set, reaching from 91 to 254, has just been obtained by a series of least-squares adjustments. The results of the first two adjustments, covering the mass range up to A = 70, have been used here to calculate the Q-values in the accompanying table.

The rotation vibration model for deformed nuclei is summarized, various matrix elements are defined, and multipole operators are evaluated. The rotation vibration Hamiltonian is diagonalized up to spin 20 and presented in a manner which is independent of the individual nucleus. Energies and wave functions for the various states in the ground state, beta and gamma rotational bands of even deformed nuclei are given in an extensive table. The use of the table is explained in detail. An extensive comparison of the rotation vibration model and the asymmetric rotator model with available experimental data indicates an approximate equivalence of the two theories in explaining the three lowest rotational bands. However, the rotation vibration model appears to have a decided advantage in explaining the few two-phonon vibrations experimentally observed. Small systematic deviations between the rotation vibration model and experiment for the energies of the vibrational bands are interpreted in terms of the Coriolis anti-pairing and blocking effects. These effects lead to lower limits on the pairing correlation energy and the critical spin above which all pairing ceases.

The theory of scattering by electric dipole interaction is developed by tensorial techniques, which permit an early separation of geometric and dynamic factors. The geometric relationships are formulated in terms of variables that represent arbitrary partial polarization of the incident and scattered gamma rays. The relevant dynamic properties of a nucleus are represented by a scalar, a vector, and a quadrupole polarizability. These polarizabilities correspond respectively to the values 0, 1, and 2 of the quantum number j which indicates the angular momentum transfer in the scattering process. The analysis of scattering according to angular momentum transfer is compared to the ordinary theory of angular distributions. The nuclear polarizability is discussed from the standpoint of different models. The magnitudes of the three polarizabilities can be determined by experiments with unpolarized nuclei but with some degree of circular polarization of the gamma rays; linear polarization contributes no additional information. Nuclear polarization is required to determine the phases of the polarizabilities.

These Q-values have been computed from masses of nuclides participating in the reaction, using a consistent set of nucleonic masses derived from all the relevant experimental data, this set, reaching from 91 to 254, has just been obtained by a series of least-squares adjustments. The results of the first two adjustments, covering the mass range up to A = 70, have been used here to calculate the Q-values in the accompanying table.

The rotation vibration model for deformed nuclei is summarized, various matrix elements are defined, and multipole operators are evaluated. The rotation vibration Hamiltonian is diagonalized up to spin 20 and presented in a manner which is independent of the individual nucleus. Energies and wave functions for the various states in the ground state, beta and gamma rotational bands of even deformed nuclei are given in an extensive table. The use of the table is explained in detail. An extensive comparison of the rotation vibration model and the asymmetric rotator model with available experimental data indicates an approximate equivalence of the two theories in explaining the three lowest rotational bands. However, the rotation vibration model appears to have a decided advantage in explaining the few two-phonon vibrations experimentally observed. Small systematic deviations between the rotation vibration model and experiment for the energies of the vibrational bands are interpreted in terms of the Coriolis anti-pairing and blocking effects. These effects lead to lower limits on the pairing correlation energy and the critical spin above which all pairing ceases.

The theory of scattering by electric dipole interaction is developed by tensorial techniques, which permit an early separation of geometric and dynamic factors. The geometric relationships are formulated in terms of variables that represent arbitrary partial polarization of the incident and scattered gamma rays. The relevant dynamic properties of a nucleus are represented by a scalar, a vector, and a quadrupole polarizability. These polarizabilities correspond respectively to the values 0, 1, and 2 of the quantum number j which indicates the angular momentum transfer in the scattering process. The analysis of scattering according to angular momentum transfer is compared to the ordinary theory of angular distributions. The nuclear polarizability is discussed from the standpoint of different models. The magnitudes of the three polarizabilities can be determined by experiments with unpolarized nuclei but with some degree of circular polarization of the gamma rays; linear polarization contributes no additional information. Nuclear polarization is required to determine the phases of the polarizabilities.
charge in the nucleus, nuclear moments, total beta disintegration energies, and gamma rays emitted by radioactive nuclei. A supplement to Directory to Nuclear Data Tabulations, 1958-61, is also included.

#608020  *2.1533
Fossan, D. B.; Herskind, B.
ROTATIONAL-STATE LIFETIMES
Physics Letters, 2:155-157
September 1, 1962
The half lives of the 2+ first excited states of W-180 and Hf-180 have been measured as (1.53±0.05)x10-9 sec. and (1.50±0.04)x10-9 sec., respectively. These results are included in a summary of a complete investigation of the enhancement of the 2+ rotational levels in nuclei, in which half lives of 2+ first rotational levels were measured for a series of even-even nuclei (150 less than A less than 190). The summary includes calculated experimental total conversion coefficients and the corresponding theoretical total conversion coefficients for comparison.

#6910041  *1.2033
Fujimura, K.
NN INELASTIC PROCESSES
Progress of Theoretical Physics (Kyoto) Supplement, 41-42:282-315
1967
Experimental data obtained from accelerators are summarized for pp inelastic scattering, and their characteristic features are shown. The data are separately compiled for the two final channels; the quasi-elastic scattering (isobar production) and the multiple production. Some cosmic ray data are also included. Twenty-five references are tabulated together with incident momentum, observed quantities and experimental methods.

#658036  *2.1533
Furnish, J.
NUCLEAR MOMENTS, A COMPILATION OF VALUES MEASURED BY SPECTROSCOPIC AND RESONANCES METHODS WHICH APPEARED IN THE LITERATURE BEFORE MAY 1964
Appendix I to Nuclear Data Sheets, National Academy of Sciences - National Research Council, Nuclear Data Project, and Brookhaven National Laboratory, Oak Ridge, Tennessee and Brookhaven, New York
May 1965
140p
Measured values of nuclear moments and of some of the auxiliary quantities from which moment values are derived are listed in this compilation. There are nine tables. The first contains the measured values for the neutron, proton and anti-proton moments. The other eight contain information obtained by paramagnetic resonance, microwave spectroscopy, quadrupole resonance, nuclear magnetic resonance, atomic and molecular beams, optical spectroscopy, optical double resonance and pumping techniques, and Mössbauer spectroscopy. Information obtained by angular correlation measurements has not been included in this compilation. Since detailed descriptions of methods of measuring nuclear moments have been presented by others only brief summaries are given with each table. In all tables the values of the magnetic moments tabulated have been adjusted to a standard value of the magnetic moment of the proton by means of the experimentally determined quantities and their adopted standard values of mu and frequency ratios. In the preparation of the compilation all data available up to May 1964 were reviewed. Unless important for the determination of the sign of a moment, older values have been omitted when superseded by more accurate results. The abbreviations used and possible adoption of new quantities, uncertainties, standard values, and corrections are explained in tabular style in the first part of the report. A section called Table of Rounded-Off Values and Index is included.

#6604066  *2.083-2.1083
Fuller, G. H.; Cohen, V. W.
NUCLEAR SPINS AND MOMENTS
Nuclear Data Sheets, A5:433-612
March 1969
A summarizing table of rounded-off nuclear-moment values is arranged by nucleus in this compilation. This summary is based on experimentally determined values of the nuclear moments which have been listed in tables according to the specific spectroscopic or resonance technique used. References are given for all values quoted. Each table is preceded by a short introduction describing the experimental technique involved and the method of calculating the moments from the measured quantities. The cutoff date for the information included is August 1968. This tabulation supersedes Nuclear Moments, Appendix I to Nuclear Data Sheets, issued May 1965.

#6904009  *3.153-2.2633
Furnish, J.
ALPHA-ALPHA ELASTIC SCATTERING: A BIBLIOGRAPHY
LA-4011-MS, Los Alamos Scientific Laboratory, Los Alamos, New Mexico
October 1968
16p
Nuclear Science Abstracts was searched from Volume 7 (1953) through Volume 22, Number 15 (August 15, 1968) for material on elastic scattering of alpha particles on alpha particles. The references are arranged numerically within each volume with the abstracts from the most recent volumes presented first.

#6909066  *2.1733
Gajewski, W.; et al.
COMPILATION OF BINDING ENERGY VALUES OF LIGHT HYPERNUCLEI
Nuclear Physics, B5:105-113
February 1967
The results of a compilation of the binding energy values of light hypernuclei (A ≤ 16) obtained by the European N Collaboration and the EFIN Nuclear Collaboration are presented. The events were analyzed by the same computer program and uniform stringent selection criteria were applied.

#6910025  *2.2533
Galloway, L. A. III; Shrader, E. F.
NEUTRON TOTAL CROSS SECTIONS MEASUREMENTS USING A "WHITE" NEUTRON SOURCE
COO-1573-6, Case Institute of Technology, Nuclear Physics Laboratory, Cleveland, Ohio
September 1966
13p
Available from CFSTI
A method for measuring neutron total cross sections using a neutron intensity spectrum continuous in energy (a "white" spectrum) and a pulsed beam time-of-flight technique is used to measure neutron total cross sections in the 2 to 10 MeV region. Total cross sections for the elements Mg, Al, Ca, V, Fe, Pd, Ag and Pb were measured to 1% average uncertainty in steps of 0.08 ns/m. Energy resolution varied from about 1.5% at 2 MeV to 3% at 10 MeV. Results of these measurements are compared with measurements on the same samples with neutrons of known energy and with measurements of other workers.
Fast neutrons from the Li^7(d,n)Be^8 reaction were used to measure the neutron total cross sections of 20 elements that have principal isotopes with magic neutron numbers and to remeasure the cross sections of other elements. Neutron energies from 3 to 15 MeV were determined by the pulsed-beam-time-of-flight technique. A Fortran program for the Hanford IBM 7090 was used to calculate the cross sections. Ramsauer structure in the twenty cross sections shows a reasonable continuity with those for the remeasured cross sections. Closed neutron shells, particularly for magic numbers greater than 28, introduce small but distinct anomalies; but closed proton shells produce anomalies only if they are combined with a closed neutron shell. Unresolved anomalies appeared in As and Sr.

These volumes are issued separately, but are cataloged as a unit. Angular distributions of neutrons and of other products of fast neutron reactions with nuclei are included. Distributions for neutron elastic scattering predominate, but distributions for inelastic scattering and products associated with them are also included. Information for the various elements is arranged in order of their atomic numbers. The comprehensive data available in October 1962 and forms two volumes: Volume I, 2 = 1 to 22 and Volume II, 2 = 23 to 94.

The neutron cross section data for the elements 2 = 21 to 40, 2 = 41 to 60, 2 = 61 to 87 are presented. The volumes are arranged strictly by chemical elements and by their isotopes. Thermal cross sections, resonance parameters, and cross section curves for one element or isotope are together on successive pages, separate from the next element or isotope. The format is fluid, varying from one page to the next, to suit the information to be presented. The reference sources are listed as close to the data as is feasible.

This paper presents a systematic derivation of nuclear cross-sections beginning with the interaction between an incident particle and the target nucleons. The replacement of the exact potential by an effective potential results, to first order, in the ordinary time-independent Schroedinger equation. This equation, including a complex and spin-orbit potential is solved for the
As a part of the evaluated nuclear data file sponsored by the Atomic Energy Commission and administered by the Brookhaven National Nuclear Cross Sections Center, this report is a compilation of the 2200 m/s neutron cross sections absorption data. The tables precede and contain data published since the Supplements to BNL-325 which were published from 1964-1966. Each element is listed separately together with all the isotopes for which the cross sections have been measured.

#6604064 *2.083-2.2783
Goldstein, H.; Reynolds, S. A.
SPECIFIC ACTIVITIES AND HALF-LIVES OF COMMON RADIONUCLIDES
Nucl. Data, A1:435-452
July 1966

A compilation is presented of the half-lives, decay constants, and specific activities of 350 well-known radionuclides with half-lives greater than one day.

#6508030 *2.2133
Goryachev, B. I.
CROSS-SECTIONS OF PHOTONUCLEAR REACTIONS TABULATED EXPERIMENTAL DATA
Atomic Energy Review, 2:71-148
1964

A summary of experimental results relating to the thresholds, cross-sections and yields of photonuclear reactions is presented. Work published in periodicals up to the middle of 1963 is tabulated and is based on a breakdown of the experimental material by elements (from sodium to the transuranium elements), the papers relating to each element being arranged in alphabetical order. Except for a few old papers, the data contained in all published works were provided, they could be adapted to the tabular arrangement employed. Two-hundred fourteen references are included.

#6608024 *2.2633
Gove, N. B.
BETA AND GAMMA TRANSITION PROBABILITIES
BNL-P-1786, Oak Ridge National Laboratory, Oak Ridge, Tennessee
1965
28p
Available from CFSTI, Conf-65118-3

A survey is presented of systematics of log ft values, half lives for gamma transitions, and hindrance and enhancement factors for E1, E2, E3, M1, M2, M3, and N4 gamma transitions. Fifteen references are provided.

#6604066 *2.083-2.1283
Gove, N. B.; Yamada, M.
SHELL-INDEPENDENT SYSTEMATICS S_n (ISOTONIC) AND S_p (ISOTOPIC)
Nuclear Data, A4: 237-263
May 1968

Graphs are presented of recent values for the separation energy of a given neutron versus mass number, S_n (isotonic), and for a given proton versus neutron number, S_p (isotopic). The graphs confirm the fact, pointed out by Yamada and Matsuno in 1961, that the separation energy of the nth proton increases monotonically (except for odd-even effects) as the neutron number is increased, showing little, if any, discontinuity at magic neutron numbers. An analogous statement can be made for the separation energy of the nth neutron as the proton number is increased. Extrapolation by means of the graphs should, therefore, be particularly reliable since it is essentially magic-number independent. Data available in the summer of 1967 have been considered.

#6908058 *2.2533
Greenwood, R. C.; Reed, J. H.
PROMPT GAMMA RAYS FROM RADIATIVE CAPTURE OF THERMAL NEUTRONS VOLUMES 1 AND 2
IIITRI-1193-53 (Volumes 1 and 2), IIT Research Institute, Technology Center, Chicago, Illinois October 1963
898p
Available from CFSTI

An atlas of thermal neutron capture gamma ray spectra measured under standard conditions using NaI(Tl) scintillation detectors, and a comprehensive compilation of the neutron capture gamma ray data which was available prior to May 1963, are contained in this report. Included are prompt gamma ray spectra, and data from 74 naturally occurring elements, together with a number of separated isotopes, unstable elements and fissionable materials. The objective of this report is to provide sufficient data concerning neutron capture gamma ray spectra to allow selection of practical uses for an analytical technique based upon the detection of neutron capture gamma rays in measurement, analysis and control technology and thereby to facilitate the use of the method in these areas.
This is a French version of the Table of Isotopes published under the direction of two Nobel Prize winners, F. Joliot and I. Curie, which is now primarily of historical interest.

The Atlas contains a table of stable isotopes, their thermal neutron capture cross-sections, and the binding energies of the neutron in the resulting nuclei. This table also includes other data needed in the analysis of gamma-spectra and the construction of gamma-transition diagrams. The main part of the book consists of gamma-spectra. Tables of gamma-ray energies and intensities taken from various sources, and gamma-transition diagrams of nuclei formed by neutron capture, are also given. There are two appendices, which contain data on conversion electrons emitted in thermal neutron capture and on gamma-rays from radioactivity which appear in gamma-spectra from the (n,γ) reaction. All information concerning gamma-rays from radiative capture of thermal neutrons published before January 1958 has been collected and systematically presented.

A compilation is presented of thermal-neutron-capture gamma-ray energy and intensity values with decay schemes, examples of original data, and fully corrected spectral distributions. An auxiliary table of data on stable isotopes with abundances, thermal-neutron radiative-capture cross sections, spins, parities, and other information of value to neutron-capture gamma-ray spectroscopy is included.

A compilation is presented of thermal-neutron-capture gamma-ray energy and intensity values with decay schemes, examples of original data, and fully corrected spectral distributions. An auxiliary table of data on stable isotopes with abundances, thermal-neutron radiative-capture cross sections, spins, parities, and other information of value to neutron-capture gamma-ray spectroscopy is included.

This table is a compilation of the atomic masses, symbols, atomic numbers, nominal weights, square roots of masses, relative abundance and absolute abundance of the isotopes of nuclides. It contains over 2400 entries.

Calculations of K-, L-, and M-shell conversion coefficients were performed for nuclei with Z from 31 to 103 taking into account the four lowest electric and magnetic multipoles. The energies of the continuum electron ranged from 1 to 1500 keV for the M-shell. The coefficients are presented as a function of the continuum electron energy, the continuum electron momentum, and the gamma-ray transition energy.

Values of the internal conversion coefficient, number of electrons per photon emitted in a nuclear transition, are presented from a new relativistic self-consistent field calculation which takes into account finite nuclear size, hole and exchange effects, experimental electron binding energies, and vacuum polarization. Coefficients are given for each value of Z; for K, L, and M electron shells and M-sub-shells and for nuclear transition multipoles E1, E2, E3, M1, M2, M3, and for various nuclear-transition energies up to 1500 keV. A program for finding values for other energies by spline interpolation is appended.

Values of particle parameters used for the analysis of angular correlations involving internal-conversion electrons are presented. Directional particle parameters are given for the K-, L-, M-sub-shells and for the total L- and M-shells. Polarization particle parameters are given for the K-shell. The calculations have been performed for the four lowest electric and magnetic multipoles for all Z-values from 30 to 103 and for electron energies varying from 6 to 1500 keV above threshold. Only the particle parameters of lower rank are given. Those of higher rank can be found from the recursion relations presented. A table of P-coefficients is included.

Coefficients for the analysis of penetration effects (nuclear-structure effects) are presented for internal-conversion coefficients and directional particle parameters. The coefficients are given for the four lowest electric and magnetic multipoles in the K-shell and for the two...
lowest electric and magnetic multipoles in the L- and M-shells. The coefficients are given for every fourth 2-value from 30 to 102 and for electron energies varying from 6 to 1500 keV above threshold. Some higher-order coefficients are given for E1 conversion. Coefficients for E0 internal conversion are included for the K- and L-shells, and values of the E0-E2 interference particle parameter are included for the K-shell.

#650080
Halperin, J.; Stoughton, R. W.
A TABULATION OF INTEGRATED NEUTRON AND FLUX INTENSITIES FOR FISSION SPECTRA FROM 0.01 TO 25 MeV
ORNL-TM-795, Oak Ridge National Laboratory, Oak Ridge, Tennessee
February 7, 1964
70p

The fission neutron spectrum was tabulated as a function of energy for several analytical expressions that were fitted to experimental data for thermal fission in U-235, U-233, Pu-239, and spontaneous fission in Cf-252. The integrated neutron density and flux were calculated for energies from 0.01 to 25 MeV in intervals of 0.05 MeV.

#660406
Hamilton, J. H.; et al.
EXPERIMENTAL VALUES OF INTERNAL-CONVERSION COEFFICIENTS OF NUCLEAR TRANSITIONS TERTIA AND K-SHELL COEFFICIENTS AND L-SUBSHELL COEFFICIENT RATIOS
Nuclear Data, A1:521-602
August 1966

A tabulation is presented of experimentally measured values of internal-conversion coefficients for the K-shell, alpha, L-shell subsheals (mostly ratios only, L1:L2:L3), and for all shells, alpha only. Results reported prior to November 1965 whose uncertainties are ± 25%, are listed and compared with theoretical values. Transition energies and spins and parities of initial and final levels, when known, are included. The theoretical values were obtained from computer interpolation, and in some cases extrapolation, of the tables of Silv and Band. A brief discussion of the methods of measuring conversion coefficients is given.

#690932
Hammond, J. P.
PHYSICAL, MECHANICAL, AND IRRADIATION PROPERTIES OF THORIUM AND THORIUM ALLOYS
ORNL-4214, Oak Ridge National Laboratory, Oak Ridge, Tennessee
April 1968
71p
Available from CFSTI

Physical and mechanical property data on thorium and thorium alloys were reviewed, and information pertinent to nuclear reactor application is compiled.

#6909070
Hardy, E. P. Jr.; Rivera, J.
HEALTH AND SAFETY LABORATORY FALLOUT PROGRAM QUARTERLY SUMMARY REPORT, MARCH 1, 1967 - JUNE 1, 1967
July 1967
16p
Available from CFSTI

This report presents current data from the HASL Fallout Program, Isotopes, Inc., Euratom Joint Nuclear Research Center, Argonne National Laboratory, and the National Radiation Laboratory in New Zealand. Radionuclide levels in stratospheric air, surface air, fallout, milk, other diet components, and tap water, are given in tabular form. The initial section consists of interpretive reports and notes covering the following topics: fallout of SNAP-9A debris; the ratio of Cs-137 to Sr-90 in monthly fallout; and strontium-90 in human vertebrae. A bibliography of recent publications related to radionuclide studies is also presented.

#6500163, #6500164
Heath, R. L.
SCINTILLATION SPECTROMETRY, GAMMA-RAY SPECTRUM CATALOGUE, SECOND EDITION, VOLUMES 1 AND 2
August 1964
Available from CFSTI

This second edition is a complete revision of the original data compilation which was issued as an AEC R and D Report (IDO-16408) in 1958. As in the original catalogue, this edition contains a collection of spectra representing the response of a scintillation spectrometer to individual radioactive nuclides. In addition to the graphs representing the response of a 3" x 3" NaI detector in a standard geometry, the data are presented in digital form for the preparation of punched-card, perforated tape, or magnetic tape libraries for data analysis. An important addition to the catalogue is data for neutron-deficient isotopes. The new edition is prepared in two loose-leaf volumes and contains data for almost 300 isotopes. All spectra are normalized to a standard set of gain scales and a text is presented which describes the fundamentals of gamma-ray spectrometry. This includes a discussion of spectrometer design, electronics, instrumental calibration, and data processing. To facilitate the use of these data, tables of detector efficiency, photopeak efficiency, and other information useful for quantitative data analysis have been included. An extensive index has also been added with separate tables of data listed according to gamma-ray energy, half-life, method of source production, and other specialized categories.

#6911011
Hemmings, P. J.; Offord, S. M.
PHOTON INTERACTION IN THE U.K.A.E.A. NUCLEAR DATA LIBRARY
AHSB(S)-R-109, United Kingdom Atomic Energy Authority, Authority Health and Safety Branch, London, England
1966
43p
Available from AEC Dep. Libraries

Photon-interaction data for 25 elements and water were collected for use in the UKAEA Nuclear Data Library. The methods of photon interaction are discussed with reference to transmission calculations. Tabulations are given of cross sections for each element, and the angular distribution for Compton scattering is given at 23 energies. The energy range considered is 0.01 to 20 MeV.

#6510012
Hillman, M.
EXTENDED TABLE OF NUCLIDIC MASSES
BNL-846(T-333), Brookhaven National Laboratory, Upton, New York
March 1964
143p
Available from CFSTI

About 7000 nuclidic masses for A = 1 to 255 were calculated based on parabolic systematics. The parabolic parameters, the masses, and energies available for emission of negatrons, positrons, neutrons, protons, and alpha particles are tabulated. The known nuclidic masses are compared with the calculated masses.
neutronics
shorter and the process total of INDIVIDUAL correction + The d
impl A2:243-262
surface The 20 are 20 The energy neutron partial VIENNA, d
a urani be
November January MARCH
Available
Vienna, THERMODYNAMIC THERMODYNAMIC plotting,
These and on Spectra transitions.
San Francisco December 1964
Available from CFSTI
Beta spectra for 397 beta−-emitters were computed at 200 points from the Fermi theory of beta decay using a correction factor for unique-forbidden transitions. These spectra, after normalization and plotting versus energy, are presented, along with experimental end-point energies, forbiddenness assignments, and half lives.

Hogan, O. H.; Zigmans, P. E.; Mackin, J. L. BETA SPECTRA II. SPECTRA OF INDIVIDUAL NEGATRON EMITTERS USNRL-TR-802, Naval Radiological Defense Laboratory, San Francisco, California December 1964 60p
Available from CFSTI

Hogan, O. L. BETA SPECTRA V. SPECTRA OF INDIVIDUAL POSITRON EMITTERS USNRL-TR-1101, Naval Radiological Defense Laboratory, San Francisco, California November 14, 1966 134p
Available from CFSTI, AD646228
Spectra for 219 positron emitters were computed, with energy intervals of .01 MeV (Nb-92 and Ta-177 at shorter intervals). The computation was based on the Fermi theory of beta decay, with a correction factor for unique-forbidden transitions. These energy spectra, after normalization and plotting, are presented, along with the data used in their computation.

Available from AEA, Vienna, Austria
This report deals with the physical, thermodynamic and transport properties of uranium dioxide and related phases of the uranium-oxygen system. It includes data and a summary of research work done in the field through 1964. The following topics are included: crystal structure, heat capacity, free energy, enthalpy, and entropy measurements, vaporization processes, surface and oxidation properties, thermal conductivity, electrical properties, optical properties, magnetic properties, and practical implications of thermodynamic and transport properties.

Honick, H. C. Pearlstein, S. (Revisor) ENDF/B SPECIFICATIONS FOR AN EVALUATED NUCLEAR DATA FILE FOR REACTOR APPLICATIONS BNL 50066 (T-467) ENDF 102, Cross Section Evaluation Center, Brookhaven National Laboratory, Upton, New York May 1966 (Revised July 1967) 114p
Available from CFSTI
This report illustrates and describes the several steps required to process measured nuclear data into a form suitable for input to a reactor design code. It also describes a second part of the ENDF system (a magnetic tape store of data), referred to as ENDF/B, and contains the specifications for a magnetic tape (or punched card) storage for evaluated nuclear data to be used for reactor design calculations. The extension of these formats to other than reactor applications has not been included in this report.

Horsley, A.; Stewart, L. EVALUATED NEUTRON CROSS SECTIONS FOR DEUTERIUM LA-5271, Los Alamos Scientific Laboratory, Los Alamos, New Mexico January 1968 113p
Available from CFSTI
This collection of tables and graphs evaluates experimental data on neutron cross sections for deuterium. Most of the experimental information has been obtained from the literature. Often these data could not be obtained in numerical form and had to be read from graphical displays. Values are recommended for neutron-deuterium cross sections in the energy range 0.0001 eV to 20 MeV to facilitate neutronics calculations.

Horsley, A. NEUTRON CROSS SECTIONS OF DEUTERIUM IN THE ENERGY RANGE 0.0001 eV TO 20 MeV Nuclear Data, A:4:321-357 July 1968
Experimental and theoretical data for the neutron cross sections of D are surveyed and values adopted for total and partial cross sections. To facilitate neutronics calculations, angular distribution functions based on n + d and the conjugate reaction p + d are given. The experimental data used, with some comments, and a brief account of the evaluation procedure are presented.

Horsley, A. NEUTRON CROSS SECTIONS OF HYDROGEN IN THE ENERGY RANGE 0.0001 eV - 20 MeV Nuclear Data, A:2:245-262 September 1966
Theoretical and experimental data for the neutron cross sections of H-1 are surveyed and values recommended for total and partial cross sections in the energy range 0.0001 eV to 20 MeV. Available experimental data are supplemented where necessary by estimates based on nuclear theory. Details of energy and angular dependence are given so that the data are complete for the purposes of neutronics calculations.

Available from AEC Ser. Libraries
Recommended values are given for neutron total and partial cross sections of H-1 in the energy range 0.0001 eV to 20 MeV. The available experimental data are supplemented where necessary by estimates based on nuclear theory. Details of energy and angular distributions are given so that the data are complete for the purposes of neutronics calculations. The recommended data in this report have been written on punched cards (3A/H data, data file number DFN 211, for the cross sections of free hydrogen atoms, and 3B/H data, data file number DFN-212, for the cross sections per hydrogen atom of a water molecule) as part of the UKAEA Nuclear Data Library.

Available from CFSTI

31
Several cross section curves are plotted for most of the elements and isotopes considered. For most of the elements and isotopes, the total cross section has been measured in the energy range under consideration. For elements with Z > 40, optical model calculations have been made for the total cross section in the energy range from 7 to 14 MeV. The calculations of Fernbach and Bjorklund were used to supplement the experimental data where there was a shortage of measured values or where an ambiguity existed between several measured values. These data are of value for historical interest.

Tables of neutron total, elastic, inelastic scattering, and absorption cross sections are presented for hydrogen, lithium, beryllium, bismuth, lead, titanium, and vanadium through 14.5 MeV. These data are of value for historical interest.

Tables of neutron cross sections for various elements at energies of 0.001 to 14.5 MeV are presented. These data are of historical importance.

Part 1, Volumes 1, 2, and 3 was issued as report UCRL-5226, revised October 1959. Part 2 was issued as report UCRL-5351, November 1958. Tables are presented of experimental differential neutron cross sections for the elastic scattering of neutrons by nuclei in the energy range of 0 to 15 MeV. Nuclear reactions induced by neutrons are also included, particularly those that are significant for reactor-type calculations. The tables include nuclei from H to Pu. These data are of historical value.

This is a description of the Experimental Cross Section Information Library (ECSIL), a system, written primarily in basic Fortran II, for the storage, retrieval, and tabular or graphical display of experimental constants associated with neutron-induced nuclear reactions.

Thresholds are tabulated for nuclear reactions initiated by neutrons, protons, deuterons, tritons, He nuclei, He+$\alpha$, and gammas (14007). Gamma-production cross sections, both integrated and differential, are tabulated. The data are categorized according to whether the gammas produced are continuum, discrete, or total over-all gamma energies (14007).

"Photon Attenuation" is a section of the Engineering Compendium on Radiation Shielding which discusses and tabulates narrow-beam attenuation coefficients and energy-absorption coefficients. The mass energy-absorption coefficient is only one of many quantities that can be used to compute the transfer of energy to the medium. The mass energy-transfer coefficient and the mass absorption coefficient are also tabulated. The principal results for attenuation coefficients and for the energy-absorption and related coefficients are listed in tables. The present compilation is based on a new analysis of all available information. Discussions of the atomic cross sections underlying the interaction coefficients are included.

The material in the first edition of BNLS-325 and its addendum, the supplement, and new data received up to May 1958 are included. Thermal cross sections, resonance parameters, and cross section curves are given.

A compilation of data on neutron cross sections and resonance parameters, published between July 1958 and November 1, 1959, is presented.
A compilation was made of the data that would be most useful in calculating the activities produced in food by high-energy electrons or gamma rays. The known isotopes of each element are listed with their atomic masses, half-life, and methods of decay. Photoneutron and photoproton cross sections are shown and both theoretical and experimental threshold values are listed. References are included for all data. The appendices include a short description of the Bremsstrahlung cross sections and a list of the common radioactive isotopes found in food irradiated by high-energy electrons. One hundred and five references are provided.

Hg-203 and Nb-95 were calibrated by a 4 pT beta-gamma coincidence method for gamma-emission rate, 2n-6S by comparison with the 1.12MeV peak of Sc-46, Na-22 by a gamma-annihilation-quanta coincidence method and by a triple coincidence method, and Sr-85 by x-gamma coincidence counting. The accuracy of the calibration in all cases was plus or minus 5%. The half life of the isomeric state of Rh-85 was measured and found to be 0.98 microseconds.

Experimental data on antinucleon-nucleon inelastic interactions published from 1961 to 1966 are summarized. Cross sections are first given for typical antiproton,proton processes at various energies. Experimental results are then summarized for the four classes of reactions: annihilation into pions; annihilation yielding K mesons; pion production without annihilation; and hyperon production. Annihilation data include cross sections for various annihilation channels, production angular distributions and mean multiplicity for produced particles, the rates of resonance production, and distribution of the production and decay angles for resonance particles. Pion production without annihilation and hyperon production data include cross sections, angular distributions, final antibaryons (baryons), and resonance production. Thirty-seven references are arranged according to the reaction types, and include the primary sources, observed quantities, and method used. Seventeen other references are given.

The compilation attempts to include the "best" cross sections of all nuclear processes involving charged and neutral projectiles with energies up to the region of about 30 MeV. No reactions with beams of neutrons, photons, or subnuclear particles are considered. Only results that are absolute or that can be normalized easily to existing absolute data are included. The first edition is limited to light-element targets up to and including fluorine, and the incident particles with a mass of one through four. It should be noted that the compilation is a graphical representation of experimental data. Tables of results that duplicate information on the curves are not given, nor has any consistent attempt been made to include theoretical information.

The reactions A1-27(n, alpha)Na-24, Mg-24(n,p), Na-24, Si-28(n,p)Al-28, Co-59(n,p)Fe-59, Co-59(n,2n)Co-58, Co-59(n, alpha)Mn-56, Ni-58(n,p)Co-58 and Ni-58(n,2n)Ni-57 were studied by an activation method from 12.5 to 21.0 MeV neutron energy. A method of calibrating well-type crystals to obtain absolute efficiency is described. The results for absolute cross sections obtained by this method for the above reactions were compared with statistical model calculations in which the nuclear transmission coefficients were derived from an average optical potential extracted from the scattering data.
scattering are included, and Legendre polynomial coefficients to describe the angular distribution of elastically scattered neutrons are given.

A review is given of experimental results on the inelastic K-N scattering at various primary momenta above 1.4 GeV/c. Reactions of the following two types are considered: (a) two-body reactions in which there are more than two particles in the final state, and (b) one-body reactions in which there are more than two particles in the intermediate and/or final state. The cross sections for each inelastic channel in the K-N scattering and its angular dependence are presented. Angular distributions of particles or resonances produced in the K-N inelastic processes and decay angular distributions of the produced resonances are shown. Polarizations of the X* particle in the K+ p → X K reaction are given. The dominant characteristics of all angular distribution in the inelastic reactions and the characteristics of all resonances produced in the K-N scattering are summarized. The 57 references are tabulated in the appendix together with the incident momentum, observed quantities, and experimental methods.

Values of some 700 half-lives of isomeric states, ranging from 10^{-12} seconds to 10 years, are presented in order of increasing half-life.

The 12 group set of photon cross sections calculated for use in STRAIT and other programs in which an isotropic scattering can be taken into account is described. All the numbers required are given in the form of SC4020 prints so that the cross sections can be used in other programs.

This table is an extension of the Table of Nuclear Spins and Moments published in 1965, and contains the nuclear magnetic and quadrupole moments known as of August 1967. Major changes made in the present revision are the omission of special values (except for identification purposes), the addition of lifetime- or field-dependent quantities or both, the omission of magnetic octupole moments, and the addition of a tabular and graphical form. The table of nuclear moment ratios (Part 3) lists magnetic moment, quadrupole moment, and g-factor ratios determined for two isotopes of the same element and for two levels of the same isotope. The experimental methods used in measuring the nuclear moment are indicated by identifying symbols which are defined in Part 1 of this table. Part 2 is the table of magnetic and quadrupole moments; Part 3, the table of nuclear moment ratios, and Part 4 contains the list of magnetic shielding constants used for each element. References for all parts of the table are given at the end.
These tables are the second installment of a list of adjusted Q-values of which the first installment for targets in the mass range 1 < A < 66 has been published recently. The least squares adjusted mass values used for the computation of the adjusted Q-values of the present table are those of a recent publication. As seen from table 2 of this paper the computation for the mass range 71 < A < 199 has been divided into eight separate adjustments.

A study has been made of the neutron cross sections and associated data for U-235. A set of data has been prepared in the ENDF/B format and includes: resolved and unresolved resonance parameters; pointwise cross sections for total and partial neutron cross sections in the energy region 100 keV to 15 MeV; angular distributions for elastically scattered neutrons; energy distributions for secondary neutrons from (n, n'), (n, 2n), (n, 3n) and fission reactions; nu, the number of neutrons produced per fission reaction; fission product yields for thermal neutron fission. A survey was made of the available experimental results for the U-235 neutron cross sections up to June 1968.

The tables contain the nuclear masses for 84 < A < 104 and 111 < A < 125 nuclear Physics, 41:330-342 1963

These tables contain data on elements and isotopes from Z = 0 to 12. Data on elements includes neutron and proton cross sections, parameters for neutron resonances, and nuclear reactions where the resulting mass is unknown. The experimental data includes the abundance or half-life and decay mode; its energy levels; all nuclear reactions resulting in that isotope; reaction cross-sections and energies.

A compilation of the available information on nuclear reactions of Al, Si, P, and S is presented.

This section of the Landolt-Bornstein tables contains data on the electron shells of molecules. The volume includes data published through September 1951. The following properties have been tabulated: band spectra of diatomic molecules, electron band spectra of multiaatomic molecules, light absorption of solutions in the infrared and visible region, ionization energies of molecules, optical rotary power of molecules, electrical and optical polarizability of molecules, magnetic moments of molecules, diamagnetic polarizability of molecules, and quanta output of photochemical reactions.

This volume contains data on the atomic physical interpretation of crystal physics as of November 1954. The following properties have been tabulated: the characteristics of symmetry, elements, the 32 crystal classes, space groups, lattice types, structures, crystal dimensions of inorganic and organic crystals, atomic and ionic radii, lattice energy, internal vibrations of crystals, electron emission, energy bonds, X-ray spectra and bonding states, electron spectra of crystals, and high frequency spectra of crystals.
This section of the Landolt-Bornstein tables contains tables on atomic nuclei and elementary particles. These tables were compiled by the authors from literature on these subjects published through November 1951. The properties tabulated are: the hyperfine structure of atomic terms and atomic lines, a table of all atomic nuclei and their properties, mass spectrographic measurements, nuclear reaction energies, electron line spectra of artificially radioactive atoms, the radioactive families, nuclear level schemes of radioactive transformations, energy schemes of the light atomic nuclei, beta-spectra, common naturally radioactive atoms and their alpha- and beta-spectra, scattering of charged particles by atomic nuclei, nuclear transition products, passage of alpha-particles, electrons and gamma-rays through matter and cosmic radiation.

This volume contains nuclear radii and other characteristic parameters of atomic nuclei which are deduced from electron scattering experiments and the observation of mesic X-rays. The first section contains data on nuclear charge distributions derived from sources other than scattering, primarily mu-mesic atoms for spherical nuclei. Data is given on energy levels and radii parameters for X-rays. The second section includes neighboring nuclides, deformed nuclei, and radii of isomers. The second section deals with nuclear radii determined by electron scattering. Both energy level and charge radii are given. Because a variety of nuclear models are available, each nucleus is discussed separately, in addition to listing the size parameters.

This first volume of the New Series of Landolt-Bornstein contains the known energy levels and transitions of all nuclei from \( A = 5 \) to \( A = 257 \). The essential parts of the material are the level and decay schemes. These are supplemented by tables and special data. The tables list the following parameters for each level: excitation energy above ground state; the total angular momentum quantum number; the parity; the isobaric spin quantum number; the half life or mean life of the state; its half-width; the known modes of decay of the states; binding energies for various nucleon combinations, and the Q-values for the relevant nuclear reactions. Much of the information is incorporated in diagrams in which the energy levels of the various nuclei are exhibited schematically together with the reactions by which they are formed. Recent references to papers dealing in a more general way with the properties of the nucleus in question are included. This compilation includes the material available through May 1960 with a few references to work done since that time.
called the F-coefficient. The GAMMA-coefficient is a slightly modified form of this F-coefficient useful for beta-spectroscopy. The coefficients mentioned above are extensively tabulated in this volume. For the jj-symbol, however, only selected ones are listed. In addition, there is a chapter dealing with angle functions. Since angular correlations are generally developed in a series of Legendre polynomials, these functions should be very useful.

The accompanying beta-decay tables present certain combinations of electron radial wave function and phase shifts for both screened and unscreened beta-decay functions. The tables for electron capture give the amplitudes and values of the electron radial wave functions at the nuclear radius for the k, l, and m shell for all values of Z (the L and M shells are excluded for very light nuclei). A brief compilation of all the relevant formulae used in the treatment of beta-decay and electron capture is given. An explanation of the details of the computations and a description of the tables is included; in addition, 56 references are given.

Brenstrahlung cross sections were calculated for screened and nonscreened targets for electrons with energies from 0.05 to 5 MeV. The cross sections were obtained by numerical integration of the Bohr-Wheeler equation for Brenstrahlung cross-sections which are differential in photon energy and in proton and electron emission angles. Values of the Brenstrahlung cross-sections which are differential in photon energy and doubly differential in photon energy and photon emission angle are tabulated.

A table is given which displays data for all radioactive and stable nuclei arranged according to atomic number with increasing mass number for each element. The criterion for the selection of data on each radioactive isotope was that of identifying it in terms of its rate and mode of decay, principal radiations, and how it is prepared. Detailed information on nuclear energy levels and transitions between these levels is given. The general policy adopted for these entries was that they be based on direct experimental information. Spin and parity assignments based wholly, or in large part, on the expectations from nuclear models were avoided. Unobserved transitions that should be present were omitted, with few exceptions. A complete bibliography of the sources of information is included.
A survey of the present state of knowledge on (n, p) and (n, 2n) reactions has been performed. The result is presented in the form of a main index to references on all elements and several special indices to different theoretical ways of approach, such as the direct-interaction concept, the continuum model (including level density theories), the discrete-level statistical model (Hauser-Feshbach), with a subsection on the optical model. There are 486 references accompanied with abstracts giving essential information pertaining to the field.

#6500074, 6500073, 6500072  *2.2453
Liikala, R. C.
UPDATED RBU BASIC LIBRARY.
VOLUME 1: RECORDS: 1-300, ISOTOPES: H-1 TO Mo-96;
VOLUME 2: RECORDS: 310-664, ISOTOPES: Mo-97 TO Au-200;
VOLUME 3: RECORDS: 665-904, ISOTOPES: Hg-196 TO Pu-240
HW-75716 (Volumes 1, 2, and 3), General Electric Company, Hanford Atomic Products Operation, Richland, Washington May 20, 1965 239p
Available from CFSTI

The revised Hanford library of cross sections, resonance parameters, inelastic spectra, and nuclear descendents (product nuclei) for neutron reactions and scattering with isotopes H-1 through Mo-96, Mo-97 through Au-200, and Hg-196 through Pu-240, is presented.

#6600820  *2.453
Liikala, R. C.
WESTCOTT PARAMETERS DERIVED FROM THE RBU BASIC LIBRARY FOR A DELTA-4 CUTOFF
Available from CFSTI

Westcott parameters were derived for the following isotopes: U-235, U-236, U-238, Pu-239, Pu-240, Pu-241, Pu-242, Am-241, Am-242, Am-243, Cm-242, Cm-245, and Cm-244. Results are presented in tabular form.

#6500069  *2.2453
Liskien, H.; Paulsen, A.
EUR-10.e, European Atomic Energy Community, Central Nuclear Measurements Bureau, Geel, Belgium November 1961 50p

Cross sections for neutron threshold reactions with Mg-24, Al-27, P-31, Fe-54, Fe-56, and Ni-58 and for the (n, alpha) reaction with Al-27 are presented. The cross section values are printed on several sheets of transparent paper enabling a survey of a given series of measurements separately as well as permitting the intercomparison of different measuring series. Uncertainties of cross section values and neutron energies are indicated. The figures of merit for the reactions are also included.

#6908011  *2.2553
Liskien, H.; Paulsen, A.
COMPILATION OF CROSS-SECTIONS FOR SOME NEUTRON INDUCED THRESHOLD REACTIONS, VOLUME 1 (A = 55), VOLUME 2 (A = 56)
EUR-119.e, European Atomic Energy Community, Central Nuclear Measurements Bureau, Geel, Belgium April 1966 100p

Cross sections are presented for neutron induced reactions in the energy range 0.1 to 20 MeV. The reactions are generally (n, alpha), (n, 2n), or (n, p). Nuclei considered are: Na-23, Mg-24, Al-27, Si-26, P-31, S-32, Ti-46, Ti-47, Fe-54, Fe-56, Ni-58, Co-59, Ni-60, Cu-63, Cu-65, and Zn-64. Comparisons on the measuring techniques and accuracies, Q-values for these and other threshold reactions, and information on the data of the various contributors are included.

#6912029, I & II  *2.2052
LRL EXPERIMENTAL CROSS SECTION BIBLIOGRAPHY, VOLUME I. BIBLIOGRAPHY, VOLUME II. CROSS SECTION DATA California University, Lawrence Radiation Laboratory, Livermore, California May 21, 1968 424p, 370p

Tables of isotopes are given showing cross sections with the energy ranges given as E-min and E-max. The number of points included in each event is noted as well as the type of reaction. The tables are arranged in three ways: 1) according to increasing A (and 2) numbers; 2) according to the reference numbers; and 3) according to the type of reaction. A separate volume is included a list of bibliographic sources, alphabetic reference list, alphabetic author list, and a numeric reference list. In all, 1,548 references are given.

#6510011  *2.1075
McCullen, J. D.; Bayman, E. B.; Zamick, L.
WAVE FUNCTIONS IN THE 1 f-, 2f-, SHELL
NYO-9901, Princeton University, Palmer Physical Laboratory, Princeton, New Jersey June 1964 75p
Available from CFSTI

Eigenvalues and eigenvectors of matrices of a two-particle interaction, calculated with respect to states of several neutrons and protons in the f-2 shell, are tabulated. The interaction is characterized by eight numbers giving the interaction energies in the eight levels available to two f-2 particles. Calculational methods and computer-prfittout results are given for Sc-43, 44, 45, 46, 47, Ti-44, 45, 46, 47, 48, 49.

#6500086  *2.2353
McGowan, F. K.; Milner, W. T.; Kim, H. J.
NUCLEAR CROSS SECTIONS FOR CHARGED-PARTICLE INDUCED REACTIONS, Mn, Fe, Co
ORNL-CFX-1, Oak Ridge National Laboratory, Charged-Particle Cross-Section Data Center, Oak Ridge, Tennessee July 1964 44sp
Available from the AEC Div. of Tech. Information

This volume is a compilation of nuclear cross section data given for charged-particle-induced reactions of Mn-55, Fe-54, 56, 57, 58, and Co-59, taken from literature before January 1, 1964. Both graphical and tabular data are included.

#6500087  *2.2353
McGowan, F. K.; Milner, W. T.; Kim, H. J.
NUCLEAR CROSS SECTIONS FOR CHARGED-PARTICLE INDUCED REACTIONS, Ni, Cu
ORNL-CFX-2, Oak Ridge National Laboratory, Charged-Particle Cross-Section Data Center, Oak Ridge, Tennessee September 1964 51lp
Available from the AEC Div. of Tech. Information

A compilation of nuclear cross sections for charged-particle-induced reactions of the nuclei Ni and Cu is presented. The compilation includes experimental cross-section data available in the literature before January 1, 1964. A number of bibliographies are included in the appendices to
This reaction list for charged-particle-induced nuclear reactions has been prepared from the journal literature for the period from 1948 through April 1960. Each published experimental paper is listed under the target nucleus in the nuclear reaction with a brief statement of the type of data in the paper. The nuclear reaction is denoted by $A(a,b)B$, where $M, Z$ (one-nucleon mass). There is no restriction on energy. Nuclear reactions involving mesons in the outgoing channel are not included.

A compilation of x-ray cross sections has been made in the range 1 keV to 1 MeV from a survey of the experimental x-ray total cross section data existing in the data files being maintained at the Lawrence Radiation Laboratory and at the X-Ray Attenuation Coefficient Information Center, National Bureau of Standards.

Preliminary graphs and tables of x-ray cross sections in the range of 1 keV to 1 MeV are given. The total cross sections presented are the sum of computed scattering cross sections and the fit to derived experimental photoelectric cross sections. Form factors available in the literature were used to compute the scattering cross sections. Where solutions to the Hartree-Fock equation were unavailable, Betwilogus values for the Thomas-Fermi model were used. L and M shell jump ratios were calculated from the theoretical values of Rakavy and Ron. The graphs for each element show the experimental data points used to obtain the fit.

Tables of $Q$-values for nuclear reactions have been calculated from the currently available mass data on the nuclides. The calculations were carried out on all stable and naturally occurring radioactive isotopes from a z of 2 to 92, as well as on a few common long-lived isotopes.
experimental data, tables of lambda-4 and lambda-6 are presented. The first factor appears in both the beta-circularly polarized gamma correlation and in the longitudinally polarized beta-gamma correlation; the second in the transversely polarized beta-gamma correlation. With the appearance of these tables, all of the Coulomb corrections found in kotani's equations are now either given explicitly, or else can be determined by simple multiplication.

#6604066 *2.083-2.2283 Marion, J. B. GAMMA-RAY CALIBRATION ENERGIES Nuclear Data, A4:301-319 May 1968

A tabulation is given of gamma-energy ranges which are suitable for calibrating high-resolution gammaray spectrometers. Data from studies of both radioactive sources and nuclear reactions are included, allowing precision calibration in the range 25 keV < E gamma < 11 MeV.


Guides to theoretical papers on nuclear-structure physics, formerly issued on cards, are presented for the years 1957-1958 and 1959-1960. The articles covered appeared in journals or reports during the given years. The bibliographies are categorized according to nuclear structure, nuclear models, nuclear properties, nuclear reactions, electromagnetic radiation, beta decay and parity, alpha decay, and special topics. Tables and computational aids as well as an author index are included.

#6604066 *2.083-2.2783 del Marmol, P. DELAYED-NEUTRON PRECURSORS Nuclear Data Tables, A6:141-151 June 1969

For delayed-neutron precursors, i.e., nuclei whose beta-decay populates neutron-emitting states of their daughters, experimental data on the following properties are compiled and evaluated: half-life, neutron-emission probability, charge and mass assignment, methods of production, beta-decay properties leading to the neutron-emitting states, neutron spectra, measured or estimated beta-disintegration energy of the parent, and neutron-separation energy of the daughter nucleus.


The tables presented here are intended as replacement of "Miscellaneous Physics Tables - Part II. Electronic Functions" (Mathematical Tables 17, National Bureau of Standards, 1941). Eight parameters have been calculated on the Standards Electronic Automatic Computer (SEAC) and are tabulated with an eight digit accuracy.

#6912007 *2.2223 der Mateosian, E.; McKeown, M. TABLE OF GAMMA-RAYS EMITTED BY RADIOACTIVE NUCLEI ARRANGED IN ORDER OF INCREASING ENERGY BNL-605, Brookhaven National Laboratory, Upton, New York May 1960 21p Available from CFSTI

Data were taken principally from two sources: the Table of Isotopes by D. Stenstrom and the Nuclear Data Sheets. To facilitate comparison of the data, all gamma rays reported in these sources were plotted on an energy scale under the names of the experimentalists responsible for the measurements. Only a very limited critical evaluation of the data was attempted.


A list of cross section data and related information has been compiled for total neutron cross sections and the reaction cross sections, (n,n'), (n, alph), and (n, 2n) in the energy region around 14 MeV. This compilation includes published data through June 1963. A gamma ray spectral atlas containing 54 sets of spectra of some of the more common elements in the Periodic Table was obtained by activation analysis with 14.8 MeV neutrons. A one and a five minute irradiation of each element was made, and spectra were recorded at various times after irradiation to emphasize decay characteristics of the radioactive products from the irradiated element. Gamma ray peaks from the reaction products were identified as well as radioactive decay products from trace impurities in each element whenever possible. Carbon and nitrogen surface contamination of several high purity samples were measured by proton irradiation. The results seem to indicate that a large percentage of this contamination is in surface layers suggesting that these contaminants may have been added during a machining process.

#6601001 *2.1153 Mattauch, J. H. E.; Thiele, W.; Wapstra, A. H. ADJUSTMENT OF RELATIVE ATOMIC MASSES Nuclear Physics, 67:73-120 May 1965

An extensive computer program has been written for, and applied to, the calculation of "best values" for the masses of atoms from measured values of mass spectroscopic doubles and those of nuclear reaction energies. The program also serves for the calculation of reaction and decay energies derived from these best values. The experimental input data are tabulated, discussed and compared with the adjusted values.


A list of alpha decay energies, binding energies of some combinations of protons and neutrons and reaction energies of various nuclear reactions for a large number of nuclides are derived by least squares adjustment of a large set of experimental data.
This is a completely new mass table derived from experimental data. Data given for each element include, in order, atomic number, mass excess, binding energy, and beta-decay energy. A list of references to the literature is included.


This handbook contains a brief discussion on the interaction of neutrons and gamma rays with matter and a discussion of various techniques used in the measurement of doses produced in the material.


The measurement of neutron flux and spectra is discussed, various methods are compared, and results of intercomparisons are given. Methods of measurement are discussed for the emission rate of radioactive neutron sources, thermal neutron flux, intermediate neutron flux, fast neutron flux, and neutron energy spectra. Neutron radiation instruments for area survey and personnel monitoring involving flux and spectrum measurements are included. Typical spectra of various neutron sources are shown.


A tabulation of experimental data, extrapolated chain yields, calculated charge distributions and a recommended value for all known fission products is presented. The fissionable nuclides covered were U-235 at thermal, fission spectrum, and 14 MeV neutron energies; Pu-239 at thermal and fission spectrum energy; and Pu-241 at thermal energy.

DEUTERON ABSORPTION COEFFICIENTS AND REACTION CROSS SECTIONS CALCULATED WITH OPTICAL MODEL POTENTIALS. Nuclear Data, A2:263-279, September 1966. Tables of deuteron absorption coefficients and reaction cross sections are presented for C, Mg, Ni, Rh, Na, and Au at deuteron energies of 4 to 28 MeV. These were obtained from optical model calculations using parameters previously published by the authors.

ANALYSIS OF THE (n, alpha) NUCLEAR REACTIONS. Energia Nucleare, 11:324-336, June 1964. In Italian

Experimental data on (n, alpha) reactions, particularly at 14 MeV, are reviewed and analyzed by statistical model formulas.


Brief details are given of the data files in the UKAEA Nuclear Data Library at 15th April 1965. The data consist principally of cross sections and related data for neutrons at energies between 0.0001 eV and 15 MeV but some photon interaction cross sections are included. A companion report currently AWRE Report No. 0-70/65 describes the conventions of the Library which stores conveniently vast amounts of data required in the preparation of input data for neutron and photon transport computer codes. The list of data files has been prepared by a computer program and can easily be updated.


This book is a guide to theoretical papers on nuclear-structure physics appearing in journals and reports during 1961 and 1962, compiled and previously appearing on index cards. The book has been made by photographing the cards themselves. The major categories for papers are: nuclear structure, nuclear reactions, electromagnetic radiation, beta decay and parity, alpha decay, tables and computational aids, and special topics. Each reference contains the following information: title, author(s), affiliation of authors, journal name, volume, page number, and year of publication. Papers that fall into more than one classification are listed under each classification. An author index is included.

This newsletter contains a summary of neutron cross section data, information and references for H, D, Li-6, Li-7, C-12, O-16, Mg, Ti, V, Mn-55, Mo Xe-133, Sm-149, Eu-151, Eu-153, Gd, Dy-164, Lu-175, Lu-177, Au-197, U-235, Pu-238, Pu-239, Pu-240, Pu-242, Cm-244, Na-23, Hf-176, Hf-177, Hf-178, Hf-179, Hf-180, Hf-174.
The Compton energy versus angle relationship and the differential and integral Klein-Nishina cross sections are presented graphically as functions of the energy and direction of the scattered photon and of the recoil electron. These graphs are intended to serve the purpose of tables. Unpolarized primary gamma rays in an energy range from 10 keV to 500 MeV are considered. The accuracy of all curves is estimated at 1 percent. The advantage of this form of presentation is the convenience and accuracy of two-way interpolation. In general, interpolated values may be obtained with an accuracy of 2 percent.

Neuert, H.; Pollehn, H.

TABLES OF CROSS SECTIONS OF NUCLEAR REACTIONS WITH NEUTRONS IN THE 14-15 MEV ENERGY RANGE

EUR-122.e, University of Hamburg, Hamburg, Germany 1963

Available from AEC Dep. Libraries

During the last few years many measurements of nuclear reactions with neutrons (n,2n), (n,p), (n,dp), (n,d), (n,He-3), (n,t), (n, alpha), and (n,n alpha) in the energy range E_n = 14-15 MeV were carried out. The tables are a compilation of the cross-sections of these reactions and provide a basis for special theoretical considerations. In some cases, they point up the need for making further investigations on more precise measurements in this field.

O'Connell, R. F.; Carroll, C. O.

INTERNAL CONVERSION COEFFICIENTS AT GAMMA-RAY THRESHOLD ENERGIES

Nuclear Data, A3:287-325

October 1967

Internal conversion coefficients, for threshold values of the gamma-ray energy, are presented for the K-shell, L-subshells, M-subshells for all values of Z in the range 6 ≤ Z ≤ 95, and for the first five electric and magnetic multipoles. These numbers are based upon the exact analytical results previously obtained by the authors for the internal conversion coefficients for any shell and for any energy for a point nucleus with no screening.

Exact analytical results (in the case of a point nucleus with no screening) for the internal conversion coefficients for any shell are presented. These results simplify considerably at
threshold values of the gamma-ray energy. Numerical results, at threshold, are obtained for the K, L(K, L, LII, LiII) and M(MII, MIII, MIV) shells for 10 values of Z in the range 5 less than or equal to Z less than or equal to 95, and for the first five electric and magnetic multipoles. The results for the K shell agree with those obtained by Spinrad. It is shown that the threshold results are actually correct to order P (momentum of electron in units of its mass). The effects of finite nuclear size are also considered.

#6500051
Otel, G. S.
Available from the Superintendent of Documents, GPO, PB 161 582

The second order term, as derived by C. Kacser (Proc. Roy. Soc. A253 (1959)), of the Born approximation series for the bremsstrahlung cross section differential with respect to photon energy, photon direction, and final electron direction averaged over initial and summed over final polarization states has been put in a form that admits to numerical evaluation for the coplanar case. The results are valid for relativistic as well as nonrelativistic incident electrons. Completely general (non-coplanar) expressions are included for the first order (Bethe-Heitler) and second order terms. However, the latter has not been given in a form amissible to numerical evaluation. Tabulated values are given for the case of the incident electron kinetic energy = 500 keV, photon energy = 450 keV, and photon direction 20° from the incident electron direction. For this case the second order term increases the differential cross section by about 50 percent for Al and about 200 percent for Au for the various angles of electron emission. A rough approximation to the integrated (over final electron direction) cross section is included and a comparison is made with experimental values. The inclusion of the second order term brings the theoretical cross section closer to the experimental values.

#6910036
Ohba, I.; Kobayashi, T.
P1-N INELASTIC PROCESSES
Proceedings of Theoretical Physics (Kyoto) Supplement, 11-12:90-195 1967

Experimental data on πp inelastic processes published between 1964 and 1966 are compiled. Attention is given to data on the quasi-two-body processes which involve the meson resonance and/or the baryon resonance. Nonresonant direct process (10 GeV/c) and strange particle production are treated separately from the quasi-two-body processes. The incident momentum, reaction, observed quantity, and method are given with each of the 122 references.

#6752001
Okada, M.
GAMMA SPECTRA OF NATURAL ELEMENTS IRRADIATED WITH FILM NEUTRONS
Radiisotopes (Tokyo), 12:234-252 May 1965
In Japanese

#6509022
Okada, M.
SHORT-LIVED NUCLIDES FORMED BY NEUTRON ACTIVATION
Nucleonics, 22:110-111 August 1964

In neutron activation analysis various types of neutron reactions are likely to occur simultaneously. To analyze an element in a sample it is necessary to consider all of these interfering reactions. A table is presented that shows the original element, the type of neutron reaction, the nuclide formed, half life, gamma energies, and other decay particles present.

#6500088
Okada, M.
22 GAMMA SPECTRA OF SHORT-LIVED NUCLIDES
Nucleonics, 21:62-63 November 1963

A supplement is presented to Data Sheet No. 47 (N5A #17-9540) that contains gamma spectra for nuclides with half lives of 0.8 sec. to 24 min.

#6912024
Otter, J. M.; et al.
EVALUATED NEUTRON CROSS SECTIONS FOR COPPER-63, COPPER-65, AND NATURAL COPPER
Available from CFSTI

Evaluated neutron cross-section data for copper-63, copper-65, and naturally occurring copper have been prepared for the ENDF/B nuclear data file. Calculations, based on nuclear systematics, were used to augment available experimental data, where required. A complete set of neutron cross-section data was prepared for each nuclide, for incident neutron energies between 10^-3 and 1.5x10^7 eV.

#6608022
Owen, G. E.; et al.
NUCLEAR REACTIONS IN LIGHT NUCLEI AND NISBIBAUSER STUDIES
The Johns Hopkins University, Baltimore, Maryland November 1965
108p
Available from CFSTI, TID22405

Angular distributions of ground and first excited state neutrons in the reaction F-19(d, n)Ne-20 have been measured at F = 0.97, 1.47, 1.97, 2.17, and 2.47 MeV at laboratory angles from 0° to 165°. Differential neutron yield curves at 0° and 120° (laboratory) were measured from 0.9 to 2.7 MeV. Integrated cross-sections yield based on the angular distributions measured in this work as well as others show peaking for the first excited state in the neighborhood of 2.0 MeV. The neutrons were detected using a single stilbene crystal spectrometer employing pulse shape discrimination. The absolute differential cross-section at 0° for the first excited state neutrons was measured at 1.97 MeV to be 7.4 ± 0.8 mb/sr, using a CA, target. The ground state distributions at energies other than 1.0 MeV do not exhibit typical stripping patterns. The first excited state distributions exhibit broad peaks around 120° (laboratory), and the 1.97, 2.17, and 2.47 MeV curves also exhibit peaking at 0°. The angular distributions are in the process of being analyzed with the plane-wave, dual mode theory of Edwards.

#6500066
Parker, K.
NEUTRON CROSS SECTIONS OF U-235 AND U-238 IN THE ENERGY RANGE 1 keV - 15 MeV, PART I: BEST CROSS-SECTIONS FOR U-238 BASED ON MICROSCOPIC EXPERIMENTAL AND THEORETICAL DATA AVAILABLE AT DECEMBER 1961
Available from AEC Dep. Libraries
Recommended values are given for neutron total and partial cross-sections of U-235 in the energy range 1 keV - 15 MeV. The available experimental data are supplemented by estimates based on nuclear theory. Details of energy and angular distributions are given so that the data are complete for the purposes of neutronics calculations. Experimental and theoretical data which have become available since the best values were determined are discussed as far as possible. 215 references are included.

#6500041 2.2463 Parker, K.
A REVIEW OF EVALUATIONS OF NEUTRON CROSS SECTIONS AVAILABLE AT SEPTEMBER 1963
EANDC(UK)-26, Atomic Weapons Research Establishment, Aldermaston, England September 1963
35p
Available from AEC Dep. Libraries

#6500076 2.2463 Pattenden, N. J.; Harvey, J. A.
TABULATION OF THE NEUTRON TOTAL CROSS SECTION OF U-235 FROM 0.07 TO 10,000 ev MEASURED WITH THE ORNL FAST CHOPPER ORNL-TM-556, Oak Ridge National Laboratory, Oak Ridge, Tennessee April 1963
35p

A tabulation of the neutron total cross section of U-235 as a function of neutron energy from 0.07 to 10,000 ev, measured with the ORNL fast chopper time-of-flight neutron spectrometer is given.

#6604066 2.083-2.2783 Pauli, H.
SHAPES OF BETA SPECTRA
Nuclear Data, A2:281-298 September 1966

A compilation of information on the measured shapes of beta spectra is given. It contains, whenever possible, the numerical coefficients describing the "normalization factor," C(W) = Nm(W)/Nh(W), where Nm(W) is the measured energy spectrum and Nh(W) is an appropriate theoretical energy spectrum. The information was collected from publications that date roughly between 1957 and July 1966.

#6908044 2.2763-3.163 Pauli, H. C.
TABLES OF INTERNAL CONVERSION COEFFICIENTS AND PARTICLE PARAMETERS
COO-1420-137, Purdue University, Department of Physics, Lafayette, Indiana August 1967
206p
Available from CFSTI

The internal conversion coefficients and particle parameters of the K and L subshells have been recalculated for the ten nuclei, 2 = 60, 54, 68, 72, 76, 80, 84, 88, 92, and 96 and ten energies 0.02, 0.05, 0.10, 0.15, 0.20, 0.30, 0.40, 0.50, 0.70, 1.00 MeV. The four lowest orders of electric and magnetic multipole transitions have been considered. The internal conversion coefficients and particle parameters of the M-subshells have been computed for the seven nuclei, 2 = 72, 76, 80, 84, 88, 92, and 96 and for ten energies = 0.02 above. The screening of the atomic electrons has been taken into account on the basis of the Thomas-Fermi-Dirac model. All static and dynamic effects of the finite nuclear size have been taken into account, assuming a sharp nuclear edge.

#6500001 2.463 Paxton, H. C.
LOS ALAMOS CRITICAL-MASS DATA
LAMS-S567, Los Alamos Scientific Laboratory, Los Alamos, New Mexico April 1964
61p

Tabulated are critical masses of simple systems, which have been measured at Los Alamos through the year 1963.

#6510014 2.463 Paxton, H. C.; et al.
CRITICAL DIMENSIONS OF SYSTEMS CONTAINING U-235, Pu-239, AND U-233
TID-7028, Los Alamos Scientific Laboratory and Oak Ridge National Laboratory, Los Alamos, New Mexico and Oak Ridge, Tennessee June 1964
145p
Available from CFSTI

A compilation of critical data obtained from experiments performed since 1945 is presented. The report supplements the Nuclear Safety Guide (TID-7016(Rev.1)) and shows the bases of the recommendations that appear in the Guide. Conversions to standard geometries is shown. Results are given for single units and multiple-unit arrays. 130 references are provided.

#6604066 2.083-2.2485 Pearlstein, S.
AN EXTENDED TABLE OF CALCULATED (n,2n) CROSS SECTIONS
Nuclear Data, A3:327-341 October 1967

The table presents, for some 500 stable and unstable targets, computed values of the (n,2n) cross section for three neutron energies in the neighborhood of 14 MeV and for a fission neutron spectrum. Cross sections for (n,3n) for fission neutrons are also given. The calculations are based on the statistical model and on empirical expressions for inelastic cross sections and level density.

#6912025 2.2563 Pennington, E. M.
ENDF/B NEUTRON CROSS SECTION DATA FOR NATURAL HELIUM
ANL-7426, (ENDF-125), Argonne National Laboratory, Argonne, Illinois October 1968
33p
Available from CFSTI

This report describes the compilation of neutron cross-section data for natural helium. The elastic scattering cross section, \( \sigma_{\text{el}} \), and inelastic scattering cross sections, \( \sigma_{\text{inc}} \), for all stable and unstable targets are included. A listing of the data in the ENDF/B format is presented, as are graphs of much of the data.
This bibliography is the fourth cumulative edit of the literature concerned with nuclear reactors, nuclear weapons, and radioisotopes which has been selected by the Radiation Shielding Information Center. It supercedes ORNL-RSC-2 and ORNL-RSC-6 (December 1964). The Key Word Index, which should serve as an aid in locating bibliographic references to a given subject, has also been revised. The pages associated with a word in the index will contain a list of documents that might be relevant to the word. This bibliography is synchronized with abstracts published as ORNL-RSC-6 in a looseleaf binder. It supercedes ORNL-RSC-2 and ORNL-RSC-6.

A table of the SU3 representations contained in a given Young diagram specifying the irreducible representations. New orbitals are constructed for all possible partitions in the 2s-1d shell.

Tables are given for the rapid identification of gamma emitters, produced by (Rgamma) and fast-neutron reactions, from their approximately known half lives and energies.

Results of new calculations and measurements of neutron resonance integrals for various isotopes are presented. Most of the integrals for various isotopes are presented. Most of the integrals involved calculations taking into consideration negative energy levels to describe more accurately low energy neutron cross sections. Ages of fission and non-fission neutrons in various mixtures, elements, and compounds are also included.

An index to experimental data on photonuclear reactions is presented. Organized by element and isotope, each entry in the index supplies quantitative information for a specific reaction on the ranges of excitation energy, source energy, detected particle energy, and emission angles covered in each reference. Information is also given on the type of measurement and the detector used.

This index, a supplement to NBS Miscellaneous Publication 277, primarily covers data published in the period dated January 1, 1965 through the middle of April 1967. Organized by element and isotope, each entry in the index supplies quantitative information for a specific reaction on the ranges of excitation energy, source energy, detected particle energy, and emission angles for reaction produced covered in each reference. Information is also given on the type of measurement and detector used.

This report describes an evaluation of Pu-240 neutron cross section data carried out for the ENDF/B file. Data were evaluated from 10^-4 to 15 MeV for the following neutron reactions: total, (n,gamma), (n,fission), (n,2n), and inelastic scattering including Legendre polynomial expansions of the angular dependence, nonelastic, and inelastic scattering including resolved levels. Graphs of the evaluated data are included in the report.

This report describes an evaluation of sodium-23 neutron cross section data carried out for the ENDF/B file. Data were evaluated from 10^-4 eV to 15 MeV for the following neutron reactions: total, elastic scattering including Legendre polynomial expansions of the angular dependence, nonelastic, inelastic including resolved levels, (n,gamma), (n,2n), (n,2n), and (n,3n). Graphs of the evaluated data are compared with experimental data in the report.

A table of alpha-emitting nuclides arranged according to alpha-energy is presented as an aid in the interpretation of alpha spectra for analytical purposes. The table includes the energy Alpha of
the emitted alpha particle, relative abundance of the alpha emission of energy Ealpha, element and mass number, half-life of the nuclide, branching fraction of alpha emission of the nuclide, and year and symbol of first author of publication with a tabulation of author symbols given at the end of the table.

#6608023 *2.2663
Pressly, R. S.
LOW ENERGY GAMMA AND X-RAY FROM RADIOACTIVE SOURCES
ORNL-P-1928, Oak Ridge National Laboratory, Oak Ridge, Tennessee
February 1966
31p
Available from CFSTI, Conf 660207-2

The methods used in the fabrication of low-energy-gamma, bremsstrahlung and X-ray gamma sources are described. The spectrums of the radiation from several sources are given with efficiency measurements. The newer radio-isotopes which are being investigated and their availability are discussed.

#6912003 *2.2563
Price, L. R.; et al.
A COMPILATION OF KN REACTIONS
UCRL-20002K\textsuperscript{\textregistered}, Lawrence Radiation Laboratory, University of California, Berkeley, California
September 1969
240p
Available from the Particle Data Group, Lawrence Radiation Laboratory, University of California, Berkeley, California

Ninety papers reporting K\textsuperscript{\textregistered} and K\textsuperscript{\textdagger} interactions. Cross section, angular distributions, and polarizations are displayed. Included are indices to the papers, as well as a complete listing of the selected K\textsuperscript{\textregistered} data, and the results of the fits to some of the data. The cutoff date for this report was July 15, 1969.

#6911026 *2.2563
Prosdocimi, A.
NEUTRON CROSS SECTIONS AND FISSION PARAMETERS OF PU-240, PU-241, AND PU-242
ENANDC(E)-74(U), European Atomic Energy Community, Central Nuclear Measurements Bureau, European-American Nuclear Data Committee, Geel, Belgium
August 25, 1960
18p
Available from AEC Dep. Libraries

A compilation of neutron resonance integrals and parameters, values for the numbers of prompt neutrons emitted in neutron and spontaneous fission, and neutron fission and total cross sections is presented for Pu-240, Pu-241, and Pu-242.

#6500634 *2.372-0.2172
RADIOACTIVITY QUANTITIES AND UNITS
ICRU REPORT 10A
October 1964
17p
Available from Superintendent of Documents, GPO

This handbook presents recommendations agreed upon at the meeting of the International Commission on Radiological Units and Measurements held in Montreux, Switzerland, in April 1962. Quantities, units, and the names of radiation are given.

#6508046 *2.382
RADIOACTIVITY - RECOMMENDATIONS OF THE INTERNATIONAL COMMISSION ON RADIOLOGICAL UNITS AND MEASUREMENTS (1962) (ICRU) REPORT 10C
November 1963
53p
Available from Superintendent of Documents, GPO

This publication contains information on the following subjects: (1) direct and relative measurements of the activity of radioactive sources; (2) low-level radioactivity in materials and its relation to radiological measurements; (3) availability of radioactivity standards - present and future requirements; (4) techniques for measuring radioactivity in samples and at low levels.

#6500149 *2.2672
THE RADIOCHEMICAL MANUAL PART 1: PHYSICAL DATA
The Radiochemical Centre, Amersham, England
January 1962
102p
Available from AEC Dep. Libraries

A table of radioisotopes which are commonly used and readily accessible is presented. The table is arranged alphabetically, and for each radioisotope the basic physical data is given. An indication of the usual method of their production and reference to the physical and chemical forms which are readily available are presented. In other tables the principal radioisotopes are classified by half-life, radiation energy, and other physical characteristics. There are also select lists of isotopes which are suitable for special purposes, such as radiography, thickness gauging, the calibration of instruments and as sources of neutrons.

#6510031 *2.2673
Rakow, A.
TABELLE ZUR IDENTIFIZIERUNG UNBEKANNTER GAMMA-SPEKTREN (TABLE FOR THE IDENTIFICATION OF UNKNOWN GAMMA SPECTRA)
Munich, Germany: Verlag Karthiweg KG, Thiemig-Taschenbucher Band 2.
1962
70p
In German

Data on gamma-emitting isotopes are tabulated, including half lives, gamma energies, and the number of quanta of each energy emitted per 100 decays.

#6912027 *2.2073
Raman, S.; Miller, J.
DATA SUMMARIES (NUCLEAR-STRUCTURE PHYSICS)
ORNL-4317-1, Oak Ridge National Laboratory, Oak Ridge, Tennessee
September 1968
212p
Available from CFSTI


#6508045,6508042 *2.082
REACTION HANDBOOK, VOLUME I: PHYSICS, VOLUME II: ENGINEERING, FIRST EDITION
February 1965, May 1965
790p, 1075p
Available from Superintendent of Documents, GPO
A condensed source of reliable data and reference information for those working in the reactor field is presented. Volume I of the Handbook is divided into two sections. Section 1, Reactor Physics and Section 2, Radiation Shielding. Section 1 contains information on reaction theory, nuclear physics, kinetic theory of neutrons, reactor statics and reactor dynamics. Section 2 gives information on sources of radiation, permissible levels of radiation, gamma-ray attenuation, neutron attenuation, geometry, ducts through shields, heat generation in shields, shield optimization with respect to weight, and shield materials. Volume 2 contains information on the following subjects: light- and heavy-water-cooled systems, liquid-metal-cooled systems, gas-cooled systems, aqueous fuel systems, liquid-metal-fuel systems, fused-salt systems, handling and control, and reactor designs. An Appendix and Index are also given.

The Reactor Handbook, Volume 3, Section 1: General Properties of Materials
AEC-3647, Atomic Energy Commission, Washington, D. C.
February 1955
610p
Available from Superintendent of Documents, GPO

This Handbook is a comprehensive and critical compilation of nuclear engineering data issued by the Atomic Energy Commission's Technical Information Service for reference use by scientists and engineers engaged in AEC reactor projects. The material presented represents the efforts of specialists in the various areas of reactor science and technology, and summarizes the accomplishments of the Commission's nuclear reactor program to date. Important nuclear reactor data are given in graphical and tabular form for the following metals and alloys: aluminum and its alloys; beryllium and its alloys; beryllia; beryllium carbide; bisilum; carbides; cements and concretes; graphite; hydrides; lithium and its alloys; magnesium, molybdenum, nickel, plutonium, thorium, titanium, uranium, vanadium, zirconium, and their alloys; cobalt-base alloys; rare earths; silicon carbide; stainless steels; tungsten; and high-cross-section materials.

Reactor Physics Constants, Second Edition
July 1963
850p
Available from Superintendent of Documents, GPO

This book of Reactor Physics Constants is a compilation of the latest and "best" values of constants, recipes, formulae, etc., which are necessary to calculate reactor characteristics. This edition is based on data available through March 15, 1961. The values presented include data associated with the fission process; selected cross section data; the statics of thermal homogeneous and heterogeneous reactors; information pertinent to the analysis of control effectiveness, long-term changes, reactor kinetics, stability, and excursions in thermal reactors; multigroup calculation methods for intermediate and fast reactors. Also included are tables of representative multigroup constants, integral fast neutron physics data with emphasis on parameters for simple source analysis, and corresponding data for more complex reactor prototypes; basic shielding constants; the constants necessary for the interpretation of counting data, and the properties of concrete, soil, and yields of both artificial and spontaneous fission neutron sources are presented in tabular form. Computer programs used in reactor physics and reactor design are described. A subject index is included.

*6908020
Reynolds, J. T.; et al.
Evaluated Cross Sections for the Hafnium Isotopes
KAPL-3527, ENDF-112, Knolls Atomic Power Laboratory, Schenectady, New York
August 17, 1967
54p
Available from CFSTI

Evaluated libraries of cross sections have been prepared for natural hafnium and its isotopes Hf-174, Hf-176, Hf-177, Hf-178, Hf-179, and Hf-180. The libraries contain total, elastic, capture, inelastic, (n,p), and (n,2n) cross sections and elastic scattering Legendre moments below 15 MeV. The most recent experimental data were used in the evaluation, and whenever data were not available, theoretical calculations were made.

*6908022
Reynolds, J. T.; Ketchen, E. E.
Strontium-90 Data Sheets
ORNL-4187, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
52p
Available from CFSTI

Data on the fuel forms of Cm-244 are presented. Properties of Cm-244 metal, Cm-203, Cm-205, CmF5, Cm-242, 242Cm203 cermets are tabulated. Forty-nine references are given.

*6908021
Reynolds, J. T.; Ketchen, E. E.
Cesium-137 Data Sheets
ORNL-4187, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
27p
Available from CFSTI

Data on Cs-137 are tabulated. Properties of the fuel forms CsCl, Cs2SO4 and cesium borosilicate glass are given. Twenty-seven references are included.

*6910039
Rose, H. J.; Brink, D. M.
Angular Distributions of Gamma Rays in Terms of Phase-Defined Reduced Matrix Elements
Reviews of Modern Physics, 39:306-347
April 1967

The theory of angular distributions of gamma-rays is developed systematically, aiming at a phase consistent derivation of angular distribution formulas for gamma rays emitted in the decay of an aligned initial state. The development starts from first principles, that is, the angular distribution formulas are derived directly from perturbation theory and all quantities introduced are carefully and explicitly defined. In particular the mixing ratios are phase consistently related to reduced matrix elements of interaction multipole operators which again are well defined in phase. Hence the
mixing ratios become physical quantities which can be extracted from angular distribution measurements and then compared in both magnitude and sign with the predictions of nuclear models (especially the independent particle model). Critical stages in the theoretical development at which either a choice of phase convention has to be made or transformation properties enter are emphasized. In the last section, reduced matrix elements of the independent particle operators are calculated explicitly for one- and two-particle states and one- and two-hole states of the independent-particle model.

Rotenberg, M.; et al. 6-j and 7-j SYMBOLS
Cambridge, Massachusetts: The Technology Press, Massachusetts Institute of Technology 1959 498p

This book contains the tabulation of the 3-j and 6-j symbols, which are the Wigner modified Clebsch-Gordan coefficients in the case of two particles and the Racah Wigner distribution coefficients in the more complicated cases of quantum mechanical problems. Their high symmetry allows a tremendous abbreviation in the length of the table, with none of the concomitant handicaps in finding a particular symbol. At most, only the sign of the 3-j symbol is changed under permutations of the parameters. A complete table is compiled in the shape of a large-scale digital computer. It probably be necessary for problems requiring symbols beyond the range that is tabulated. The 3-j and 6-j coefficients are tabulated in steps of 1/2 or any angular momentum whose magnitude is less than or equal to 8 units. The 3-j symbol with all magnetic quantum numbers zero is tabulated for \( \frac{1}{2} \leq 16 \) units. A complete listing of formulas involving the 3-j and 6-j symbols, with brief mention of the 8-j and 12-j symbols.


This compilation presents existing data on the differential cross sections for the reactions antiproton-proton into ten different antihyperon-hyperon channels.


A study of the fast critical reactor 2PR-III Assembly 48 has been performed using ENDF/B cross sections. Minor alterations of the ENDF/B data were made to conform with the format requirements of the cross section preparation programs. A comparison was made of experimental and calculated results for various integral quantities, including central reactivity warchs, fission and capture ratios, and Doppler and sodium-void coefficients. The comparison indicated some possible areas of uncertainty in the basic cross section data. Real and adjoint neutron spectra were also computed for comparison with calculated and measured results. The study was performed primarily to test the unmodified ENDF/B data file in cooperation with the Cross Section Evaluation Working Group's evaluation program. For this reason modifications to the ENDF/B data were limited to minor alterations and energy range corrections.

Sasse, S. TABLES OF TOTAL GAMMA ATTENUATION COEFFICIENTS OF THE ELEMENTS AT 0.01 MeV TO 100 MeV AROTH-1040, Technische Hochschule, Arbeitsgruppe füre Bautechnischen Strahlenschutz, Amersfoort, Nederland July 1967 28p Available from AEC Dep. Libraries

The total attenuation coefficients were extrapolated from the values for 12 elements experimentally determined by Hubbell and Berger. The results are tabulated.

Schintilemeister, J. DER ISOSPIN VON ATOMKERNE (THE ISOSPIN OF ATOMIC NUCLEI) Berlin: Akademie-Verlag 1960 163p

In German

Isobaric spin and similar states of atomic nuclei, isobaric spin and the hypothesis of the density independence of nuclear forces, and the isotopic spin of light nuclei are discussed. Topics covered include comparison of the forces which act between nucleons, the concept of isobaric spins, selection rules, level displacement in isobaric nuclei on account of the Coulomb energy and the mass difference between the neutron and proton, isobaric spin of several nuclei, ambiguity of isobaric spins, isobaric spin and gamma radiation and effects on nuclear excitation, isobaric spin and photodisintegration of the types (gamma,alphas) and (gamma,sc), isobaric spin of mesons and light particles, the quantum mechanical formulation of the density independence, energy levels of isobaric nuclei, selection rules for isobaric spin and nuclear reactions, the isotopic spin of nucleons, and levels of light nuclei.

Schirmer, W.; Waeckerl, N. TABLE OF SPECIFIC ACTIVITIES OF THE NUCLIDES WITH \( n = 88 \) TO \( Z = 194 \) Actinide Reviews, 1:125-134 1968

The specific activities of all the nuclides of the elements from Ra through KU were calculated on the basis of the decay data published by A. H. Wapstra (Actinide Reviews, 1:39-53, 1967). Data for short-lived nuclides included mainly in view of their genetic relation with longer-lived nuclides.
#6608014
A TABLE OF BINDING ENERGIES AND PARTICLE SEPARATION ENERGIES FROM A MASS LAW
LA-3380-MS Los Alamos Scientific Laboratory, Los Alamos, New Mexico
September 1, 1965
108p
Available from CFSTI

A table of approximately 10,000 masses for nuclei with 22 ≤ A ≤ 120 and 22 ≤ N ≤ 184 is presented. The mass law used is one developed for application to the astrophysical problem of nucleosynthesis by rapid neutron capture.

#6604066
Seth, K. K.
S-WAVE NEUTRON STRENGTH FUNCTIONS
Nuclear Data, A:299-332
September 1966

The concept of neutron strength functions and the experimental methods of measuring them are reviewed briefly. S-wave neutron strength functions, S(0), obtained from experimental data on neutron total cross sections at energies below 700 keV are presented in tabular form in collaboration with private communications published between 1954 and July 1966 have been covered. Recommended values of S(0) are listed and plotted as a function of atomic weight. Theoretical predictions based on the nuclear optical model are also presented for purposes of comparison.

#6910009
Shirley, D. A.
TABLE OF HYPERFINE FIELDS
New York: John Wiley & Sons, Inc. 1968
6p

Hyperfine fields are tabulated for impurities in Fe, Co, and Ni; a few entries are also given for Gd as host. References are not exhaustive. Columns 2 and 3 are, respectively, the impurity and host elements. Column 3 gives the isotope number. Column 4 gives the solute concentration in atomic percent. The fields appear in Column 7. No hyperfine-anomaly corrections have been made.

#6910010
Shirley, V. S.
TABLE OF NUCLEAR MOMENTS
New York: John Wiley & Sons, Inc. 1968
57p

Small differences between the mean squared charge radii of excited states and ground states of nuclei have been deduced from isomer shifts of recoilless gamma-rays. The results of such experiments to date are summarized in this table.

#6510009, 6510003
Singsu, R. M. Jefferen, R.
TABLES FOR THE ANALYSIS OF BETA-GAMMA ANGULAR CORRELATION DATA INVOLVING FIRST-FORBIDDEN BETA-TRANSITIONS IN A 1^<=(B) + 2^<=(Y) + 0^<=CASCADE
TID-1974, Purdue University, Department of Physics, Lafayette, Indiana
August 1963
62p
Available from CFSTI

Formulas are presented for the shape factors, anisotropy coefficients, gamma circular polarizations, and parameters representing the contributions of the various nuclear matrix elements for 1^<=(B) + 2^<=(Y) + 0^<= transitions. The first three quantities are tabulated for a variety of parameters and energies. Volume I contains values for Tm-170, Re-186, and Ir-194; values for Pr-144 and Eu-152 are tabulated in Volume II.

#660406
Skrzka, J.J.; Hertel, J.; Hetz-Schmidt, T.N.
COMPILED OF ELECTROMAGNETIC TRANSITION RATES IN LIGHT NUCLEI (A ≤ 40)
Nuclear Data, A2:347-401
November 1966

Data on absolute electromagnetic transition rates in nuclei of the lp- and fs-ld-shells are presented. Table I lists the experimental results for partial widths with quoted errors, transition strengths, and reduced transition probabilities. This table also contains averages and mean errors of the averages in cases where more than one experimental result on a certain transition is available. Table II contains additional information, mainly mixing ratios which were used to calculate transition strengths in mixed transitions, as well as widths or lifetimes which could not be further reduced. Branching ratios are not listed. Table III presents a survey of mean transition strengths and of standard deviations from the mean values for the various multiple types of transitions, subdivided according to the isospin selection rules. The distributions of the transition strengths are presented in a number of histograms. The dependence of the transition strengths on the mass number and excitation energy, as well as T-forbiddenness factors, are discussed and in some cases graphically illustrated.

#6500022
Smith, D.R. (Editor)
CHARGED PARTICLE CROSS SECTIONS - NEON TO CHROMIUM LA-2424, Los Alamos Scientific Laboratory, Los Alamos, New Mexico
June 1960
117p
Available from CFSTI

Plots of cross sections for charged particle reactions are presented for Ne-20 through Cr-53. A bibliography is included.

#6908052
Smith, J.R.; Grimesey, R.A.
AN EVALUATION AND COMPILED OF NEPTUNIUM-237 CROSS SECTION DATA FOR THE ENDFB/B FILE
IN-1182, Idaho Nuclear Corporation, Idaho Falls, Idaho
May 1969
85p
Available from CFSTI

This evaluation of Np-237 neutron cross-section data was undertaken as part of the effort by the Cross Section Evaluation Working Group (CSEM) to assemble the first version of the Evaluated Nuclear Data File B (ENDF/B). Graphs of calculated and experimental data from which the various files for this isotope were constructed are included along with complete documentation over the energy range from 0.00001 eV to 15 MeV.

#6604066
Sood, D.C.
COMPARATIVE STUDY OF TWO-PARAMETER MODELS FOR PREDICTING ROTATIONAL ENERGIES IN EVEN-EVEN NUCLEI
Nuclear Data, A4:281-300
May 1968

An objective comparison of the various two-parameter models proposed for the description of rotational states in even-even nuclei is carried...
out to assess the reliability of their extrapolation or prediction of as yet unobserved levels. The comparison involving energy ratios $E(1)/E(2)$ contains no arbitrarily adjustable parameters. The semi-empirical formula

$$E(1) = A \left[ 1 - \frac{(B/A)(I+1)}{I(N(B/A)(I+1))} \right] I(I+1)$$

is seen to give the best fit. A least-squares fit to observed levels with this formula gives a rms deviation of better than 0.5% in nearly all cases. The calculated values of the rotational constants $A$ and $B$, and the predicted energies of rotational levels with spin up to $I = 16$ are tabulated.

#6509018 *2.473 Spinrad, B.I.; Meier, R.W.

The activities of the committee are summarized including comments concerning the status and accomplishments of the committee at the end of the period. Information is included on committee initial formation and functions and organization. The committee was concerned with heavy water lattice physics, lattice evaluation, thermal reactor neutron spectra, thermal reactor burnup physics, criticality predictions, irradiation experiments, nuclear data, computer codes, detector standardization, and critical experiments.

#650083 *2.2683 Stegun, I.A. (Editor)

The primary purpose of these tables is to assist in the theoretical analysis of experimental data on beta-ray spectra. This analysis involves the use of the gamma function of a complex argument, a function which cannot be readily evaluated and which has not yet been extensively tabulated. It is hoped that the present tables can be effectively utilized for most of the current and of the foreseeable needs of analysis of beta-spectra.

#6508059, 6604018 *2.2573 Stehn, J.R.; et al.
NEUTRON CROSS SECTIONS VOLUME I, 2 = 1 to 20, VOLUME III, 2 = 98 to 98

The neutron cross section data for the elements $Z = 1$ to 20 and $Z = 98$ to 98 are presented. The volumes are arranged strictly by chemical elements and by their isotopes. Thermal cross sections, resonance parameters, and cross section curves for one element or isotope are together on successive pages, separate from the next element or isotope. The format is fluid, varying from one page to the next to suit the information to be presented. The reference sources are listed as close to the data as is feasible.

#6510013 *2.373 Steigelman, W.H.
RADIOISOTOPE SHIELDING DESIGN MANUAL NYO-10721, Franklin Institute, Laboratories for Research and Development, Philadelphia, Pennsylvania July 1963 135p Available from CFSTI

Forty-two radioisotope sources, 5 additional equilibrium combinations of parent-daughter isotopes, and mixed fission products with 5 decay times are considered. Graphs are given of unshielded dose rate vs. distance from a point source, for gamma, $X$, and neutron radiations. Dose attenuation vs. shielding thickness is plotted for 5 shielding materials - water, concrete, steel, lead, and uranium. Methods of calculation are described.

#6608021 *2.2673 Stelson, P.H.
INFORMATION ON CONVERSION COEFFICIENTS FROM COULOMB EXCITATION AND LIFE-TIME MEASUREMENTS ORNL-P-1419, Oak Ridge National Laboratory, Oak Ridge, Tennessee July 1965 22p Available from CFSTI, Conf-650525-5

This is a paper presented on the International Conference on the Internal Conversion Processes, Vanderbilt University, Nashville, Tennessee, May 10-13, 1965. It reviews the available information on total E2 conversion coefficients for even-even $Z^2 + 0$ transitions for rare-earth and very heavy nuclei. Also, tables of conversion coefficient data, both experimental and theoretical, are presented.

#6604066 *2.083-2.1483 Stelson, P.H.; Grodzins, L.
NUCLEAR TRANSITION PROBABILITY, B(E2), FOR O(Ng^2) FIRST TRANSITIONS AND DEFORMATION PARAMETER, beta,
Nuclear Data, A1:21-102 December 1965

Adopted values for the reduced electromagnetic transition probability, $B(E2)$, from the ground to the first-excited $2^+$ state of even-even nuclei are given in Table I. Values of beta, the quadrupole deformation parameter, and of tau, the mean life of the $2^+$ state, are also listed there. Table II presents the data on which Table I is based, namely the experimental results for $B(E2)$, values with quoted errors. The literature has been covered to January 1965. The ratio of the value of beta to values expected from the single-particle model is presented graphically as a function of neutron number. As an indication of the extent of collective features in nuclear structure, this ratio seems a better measure than the value of beta itself.

#6907012 *2.2573 Stephenson, T.; Prince, A.; Pearlstein, S.
CROSS SECTION EVALUATION CENTER; EVALUATION OF THE NEUTRON CROSS SECTION OF MANGANESE FOR THE ENDF/B LIBRARY BNL-50060(T-463), ENDF 101, Brookhaven National Laboratory, Upton, New York June 1967 54p Available from CFSTI

This report describes the collection and choice of Mn-55 data placed in the proper format for the Evaluated Nuclear Data File - Version B, ENDF/B. These data are subject to further examination and possible revision and do not necessarily constitute the data for Mn-55 which are contained in the current ENDF/B Library, although any changes that occur should be minor. The description of how the
evaluation was performed is contained in Sections III and IV. Section III deals with the manganese cross section below 80 keV, and Section IV deals with the manganese cross section from 80 keV to 20 MeV. A complete listing of the Mn-55 data in the ENDF/B format is presented at the end of this report.

#6908009  *2.2573*
Stewart, L.
EVALUATED NEUTRON CROSS SECTIONS FOR TRITIUM
LA-1150, Los Alamos Scientific Laboratory, Los Alamos, New Mexico
April 1965
75p
Available from CFSTI

Neutron cross sections on T are tabulated from 0.025 eV to 20 MeV. Graphs and tables are presented of the total, elastic, nonelastic, (n, gamma), (n, 2n), and (n, 3n) excitation functions, as well as for the angular distributions of the neutron elastic scattering from T.

#6908024  *2.2273*
Storm, E.; Israel, H. I.
PHOTON CROSS SECTIONS FROM 0.001 TO 100 MeV FOR ELEMENTS 1 THROUGH 100
LA-3753, Los Alamos Scientific Laboratory, Los Alamos, New Mexico
June 1967
297p
Available from CFSTI

The report contains tables of photon cross sections in the range 0.001 to 100 MeV for 2 ≤ Z ≤ 100. Supplementary tables of mass attenuation coefficients for air, water, concrete, and sodium iodide; x-ray atomic energy levels; line energies; average energies; relative intensities; and fluorescence yields are also presented.

#6509006  *2.2473*
Strain, J. E.; Ross, W. J.
14 MeV NEUTRON REACTIONS
ORNL-3672, Oak Ridge National Laboratory, Oak Ridge, Tennessee
January 1965
50p
Available from CFSTI

Reactions observed when pure, naturally occurring elements (2 ≤ Z ≤ 20) are irradiated with unmoderated 14 MeV neutrons produced by the T-3(d, n)He-4 reaction are listed. Cross sections are calculated where possible, and normalized standard gamma spectra are collected. These standard spectra are equivalent to the gamma radioactivity that will be produced when 1 g of the pure element is irradiated for a given length of time at a 14 MeV neutron flux of 5 × 10^10 neutrons per sq. cm per sec. If more than one radionuclide is produced, a spectral decay is presented. These spectra may be used to predict accurately the sensitivity of nondestructive spectral analysis of almost any sample.

#6500085  *2.175-2, 2673*
Strominger, B.; Hollander, J. M.; Seaborg, G. T.
TABLE OF ISOTOPES
Reviews of Modern Physics, 30:585-904
April 1958

The table represents a complete list of all the radioactive and stable isotopes of the elements, together with a number of their salient features, as recorded in the literature or by private communications as of approximately February 1958. The system of references developed for the last edition of the Table of Isotopes has been retained. The format of the table is as follows: The first column contains the atomic numbers, chemical symbols, and mass numbers of the nuclear species. An individual entry has been made for each nuclear state whose half-life is long enough for the species to have been separated chemically or mechanically from other activities; the shortest half-life so listed is approximately 0.01 sec. Information about metastable states with half-lives shorter than 0.01 sec. is included with the other gamma-ray transition data. An exception to this policy is made when the mass assignment of a short-lived metastable state is not known; in this case a separate entry is made for the activity regardless of the half-life. Metastable excited states listed in this column are denoted by the superscript "m" following the mass number and in those cases in which two or more excited states are listed, they are distinguished by use of superscripts "m1" and "m2". Half-life values are listed without qualification where the determination has been a direct measurement of decay rate. The half-lives of nuclear states not included as separate entries are included in column four. An attempt has been made to list the most precise value or values first. The entry denoting mode of decay is preceded by the special symbol for radiation. The degree of certainty of each isotopic assignment is indicated by a letter code. The means by which the mass assignments were made are tabulated. Data on genetic relationships are located in column three under the class-assignment information. The relative isotopic abundances for the elements are given in accordance with the atomic mass values listed in the report by K.T. Bainbridge and A.O. Nier, except as supplemented by more recent values. Unless otherwise stated, the measured value of the nuclear moment is that of the ground state. The energies of particles and photons are included together in column four. Energy-level diagrams have been drawn in many cases; these are not necessarily complete representations of the data, but include only those features which seem to be reasonably well established and unambiguous.

#6910037  *1.2073*
Sumi, Y.
MESON-NUCLEON ELASTIC AND CHARGE EXCHANGE PROCESSES
Progress of Theoretical Physics (Kyoto), Supplement, 41-2:3-31, 1967

An extensive survey of experimental data on elastic pN and KN interactions up to 30 GeV/c is systematically arranged and presented. The data include total and total elastic cross sections as a function of the incident laboratory momentum (p), elastic differential cross sections, polarization of recoil proton, and charge-exchange scattering. Values of p, observed quantities, and the experimental technique are presented for the 113 references, which represent the main part of the experimental information published from 1960 to 1966.

#6910038  *1.2073*
Sumi, Y.; Yoshida, T.
NUCLEON- AND ANTIMUON-NUCLEON ELASTIC AND CHARGE EXCHANGE PROCESSES
Progress of Theoretical Physics (Kyoto), Supplement, 41-42:53-89, 1967

Experimental data on nucleon-nucleon and nucleon-antinucleon elastic interactions in the high energy region are summarized and presented. The data include total and total elastic cross sections; a singular distribution of elastic and charge-exchange cross section; and recoil proton polarization, together with the ratio of the real to imaginary forward amplitude in the pp elastic scattering. In the appendix several kinematical quantities are tabulated. Values of the incident laboratory momentum, observed quantities, and the methods used are presented for the 67 references which cover a main part of the experimental information published from 1960 to 1966.

51
A compilation is presented of data on the thermodynamic properties of materials of interest for nuclear technology. The data were released in the Symposium on the Thermodynamics of Nuclear Materials, which was held in Vienna, May 1965. The properties covered include: heats of formation, free energies of formation, entropies of formation, and standard entropies at 298° and at other temperatures; enthalpy increments; free energies; entropy increments; enthalpies, free energies, entropies of transitions; heats, free energies, entropies of reaction; heat capacities; vapor pressures; thermodynamic properties of mixtures; melting points, densities; surface tensions; and lattice parameters.

Graphs are presented of penetration factors, Pn, to enable neutron spectroscopic factors, S, to be evaluated directly from proton elastic and inelastic-scattering isobaric-analog-resonance partial widths, \( \gamma_n \), according to the relation

\[
S = (N + 1-Z) \gamma_n \gamma_n P_n,
\]

where N and Z are the numbers of target neutrons and protons, respectively. Also presented are graphs of the resonance mixing phase, \( \phi \), which is required in the extraction of \( \gamma_n \) from data. The Z-range is 16 to 82 in varying steps convenient for interpolation. The angular momentum range is \( l \leq 10 \) for \( J = 1/2 \) \( (J > 0) \). A standard set of optical-model parameters is used.

The implications of the new determination of e/h using the ac Josephson effect in superconductors for both quantum electrodynamics (QED) and our knowledge of the fundamental physical constants are analyzed in detail. The implications for QED are investigated by first deriving a value of the fine structure constant alpha from experimental input data, which do not require the use of QED theory for their analysis. These include the Josephson-effect value of e/h, the Faraday constant, the gyromagnetic ratio of the proton, the magnetic moment of the neutron, the ratio of the ampere as maintained by the United States National Bureau of Standards to the absolute ampere, and certain accurately known auxiliary constants.

The International Atomic Energy Agency initiated a project to assist in disseminating and assessing data on important nuclear materials. Thus, the symposium was organized as an attempt to present and appraise the results of the work on thermodynamics of nuclear materials; to assess their reliability and further to discuss the application of the results and principles of thermodynamics to the solution of current problems in nuclear technology. Particular emphasis was placed on the thermodynamics of actinides and vaporization processes.
elementary particles interacting with a proton. The particles $\Sigma^{+}$, $\Sigma^{0}$, $\Sigma^{-}$, $\Lambda$, and $\Xi$ were not plotted, being so close in mass to one of the particles that has been included that they cannot be distinguished graphically. Mass, energy, and momentum are expressed in terms of energy.

*2.2573

Trubey, D.K.; Emmett, M.B. 
A COMPARISON OF FIRST- AND LAST-FLIGHT EXPECTATION VALUES USED IN AN 0SR MONTE CARLO CALCULATION OF NEUTRON DISTRIBUTIONS IN WATER 
ORNL-635, Oak Ridge National Laboratory, Oak Ridge, Tennessee 
1965 
25p

Available from the RSIC, ORNL, Oak Ridge, Tennessee 

The distributions in water of fast neutrons from isotropic fission and 14 MeV point sources have been calculated with OSR code. Two methods of scoring the same case histories were used to estimate the energy distribution and the dose rate. The estimators employed were the "first-" and "last-" flight expectation, or statistical estimation, values. The first-flight estimator was found to be somewhat better due to the importance of the first flight in the total transport of the neutrons. The results are in good agreement with RENUPAX moments method results and with experimental results.

*2.2283

Uken, E.A.; Steele, T.W. 
TABLE OF THE ISOMERIC STATES OF STABLE NUCLIDES 
NIM-41, National Institute for Metallurgy, Johannesburg, South Africa 
June 15, 1966 
29p

Available from AEC Dep. Libraries 

This set of tables lists all known isomeric transitions of the stable nuclei, obtained by the nuclear reactions marked with a cross in "Table of Neutron Cross Sections." The reacting stable nuclide and its isotopic abundance in nature (quoted as percentages) are cited in increasing order of atomic number. Opposite each reaction are listed the half-life and the gamma-ray energies observed for the transition of the isomeric state as well as of the ground state. Gamma-ray energies are shown in order of decreasing intensity. In the last column the calculated Q-value, in MeV, is cited to indicate the energy balance of the reaction.

*8.182-2.782

URANIUM-CARBON AND PLUTONIUM-CARBON SYSTEMS, A THERMOCHEMICAL ASSESSMENT, A REPORT FROM THE PANEL ON THERMODYNAMIC PROPERTIES OF THE URANIUM-CARBON AND PLUTONIUM-CARBON SYSTEMS HELD IN VIENNA, 8-12 OCTOBER 1963 
Vienna, Austria: International Atomic Energy Agency, STI/DOC/10/14 
February 1963 
48p

Available from International Atomic Energy Agency 

Thermodynamic data reported by various laboratories on U-C and Pu-C systems are reported. Efforts were made to resolve various discrepancies. Sections are included on preparation of uranium carbides, phase diagrams, heat capacities and thermal properties of the actinide carbides, heat of formation of the uranium carbides, vaporization studies, high-temperature equilibrium measurements, and the Pu-C system.

*2.2583

Vidal, R.; Roullier, F. 
TABLE OF RESONANCE INTEGRALS 
Preprint No. CH-23/75, International Atomic Energy Agency, Vienna, Austria 
1966 
14p

Available from the International Atomic Energy Agency 

In French 

Tables of resonance integrals are presented which have been reduced to a common definition using $\Sigma_{0} (b-10) = 58406, \Sigma_{0} (Au) = 98.86, E_{r} = 0.0256eV, I(Au) = 15406$ as standard values and corrected to a lower cutoff of $E_{r} = 0.55eV$. The corrected resonance integrals are tabulated for several elements, references are given for each measured value and an adopted value recommended. A table of adopted resonance integrals for the natural elements and for several isotopes is also presented.
Graphs are presented of the binding energy of two neutrons as a function of $N$, of two protons as a function of $Z$, and of alpha-particle decay energy as a function of $N$. The range covered is $Z = 10$ to 102 and $N = 10$ to 154. Values shown are based on a 1964 mass adjustment.

Basic formulas and tabulations of numerical coefficients are provided for the analysis of data on angular correlations of successive radiative transitions from aligned nuclei. The formulas are extracted from an earlier published development of the angular-correlation formalism. This development is based upon a factored version of the equations which is used to extend the application from the usual two-step cascade to general multiplet cascades. The resulting flexibility and greater efficiency of the formalism are utilized to provide an extensive tabulation of coefficients for a variety of specific applications. Coefficients are provided for analysis in terms of either the population parameters or the statistical tensors of the aligned state. The coefficients can be used for the thermal neutrons, alpha-particle or linear polarizations, and triple correlations involving one or more intermediate unobserved radiations. In contrast to previous tabulations of coefficients for single transitions, all A-chains of the present tables have been extended to include multipolarities through octupole. The overall length of the tables, however, remains about the same as previous tables because of the greater efficiency of the factored formalism. The range of spins covered by the tables are $J \leq 11/2$ for the initial aligned state and $J \leq 21/2$ for the final state of cascade.

Nuclear information is presented which has been systematized according to some nuclear property, such as isotopic abundance, or some special topic such as neutron strength function. Ten tables are included which are self sufficient; explanations of abbreviations used and policies followed in each compilation are given at the beginning, and all relevant references at the end of that compilation.

A collection of diagrams showing positions and properties of nuclear energy levels, characteristics of radioactive decay and nuclear reactions, together with a tabular compilation of the experimental data and bibliographic references to the original papers is presented.
Brief descriptions of comprehensive tables of Clebsch-Gordan, Racah, Wigner, and related coefficients are presented in the first section and, in the second, of tables of coefficients of Legendre polynomials in angular-distribution expressions. The present lists, which cover material available to us up to June 1966, supersede those in Section 3 of the 1958 Directory to Nuclear Data Tabulations and its supplements. Bibliographies of some abridged tables and review articles in the same field are included in the third and fourth sections.

#6509011
Weaver, L.E.; Strom, P.O.; Killen, P.A.
ESTIMATED TOTAL CHAIN AND INDEPENDENT FISSION YIELDS FOR SEVERAL NEUTRON-INDUCED FISSION PROCESSES
USNRDL-TR-633, Naval Radiological Defense Laboratory, San Francisco, California March 5, 1963
46p
Available from CFSTI

Calculated estimates are presented of unmeasured total chain yields and of independent fission yields for the fission products resulting from fission-spectrum and 14 MeV neutron bombardment of U-233, U-235, U-238, and Pu-239 and from thermal neutron bombardment of U-233, U-235, and Pu-239.

#6908037
Kescott, C.H.; et al.
A SURVEY OF VALUES OF THE 2200 m/s CONSTANTS FOR FISSILE NUCLIDES
Atomic Energy Review, 3:5-60 1965

A study of the most probable values of the 2200 m/s constants for the fissile nuclides U-233, U-235, Pu-239, and Pu-241 has been undertaken, using a least-squares fitting method. The various experimental data have been carefully reviewed. A detailed discussion of the problems involved, especially in the accuracy of the individual measurements and of the final output values, is given. The errors shown are quoted as standard deviations.

#6711019
Wilson, B.S. (Editor)
Amersham, England: Radiochemical Centre 1966
327p

The manual is intended as a guide for all who use radioactive substances professionally; its aim is to help the individual user choose the right material for his purposes and to make effective use of it. The text portion of the book includes information on preparation of radioisotopes; radiation sources; problems of using radioactive materials; radioactive chemicals as pharmaceuticals: quantification of radioactive substances; radiological safety and waste disposal; packaging and carrying radioactive materials; ownership and use regulations and a guide to literature. The tabulated data includes: physical data on the more common radioisotopes, including physical characteristics, availability, beta and gamma ray energies and half-lives; radiation sources, synthetic routes to labelled compounds, decomposition rates of labelled compounds, measurement and safety.

#6509040
Wing, J.
A COMPARISON OF NUCLIDIC MASS EQUATIONS WITH EXPERIMENTAL DATA
ANL-6814, Argonne National Laboratory, Argonne, Illinois January 1964
8p
Available from CFSTI

Existing mathematical expressions for the systematics of the nuclidic masses, binding energies, and nuclear separation energies are summarized. The calculated values of the nuclidic mass equations of Fermi, Roman, Bussetti, Cameron, Tsen, Fidell, Seeger, Ayres et al., Kummel et al., Thieberger et al., Zeldes, Green, Levy, Baker, and Wing-Fong (a new mass equation) are compared with the available experimental data. Improvements are suggested for some of the observed deviations in these equations. Some expressions for the isobaric parameters $B_A$ and $A$ (nuclear charge of the most stable isobar), and for the pairing and shell effects are compared with the experimental evidence. An improved empirical formula for $A$ and empirical equations for the deformation parameter beta are presented.

#6509017
Wing, J.
ISOMERIC-YIELD RATIOS IN NUCLEAR REACTIONS (WITH A COMPIILATION OF EXPERIMENTAL DATA)
ANL-6598, Argonne National Laboratory, Argonne, Illinois September 1962
145p
Available from CFSTI

Experimental procedures are briefly described for the determination of the ratios of cross sections for the formation of isomeric states in nuclear reactions. Much available experimental data on isomeric yield ratios in nuclear reactions is compiled. Some of the collected data are discussed in terms of angular-momentum effects in nuclear reactions.

#6908042
Wittkopf, W.A.; Roy, D.H.; Livolsi, A.Z.
108p
Available from CFSTI

As part of the cooperative effort of the Cross Section Evaluation Working Group organized at Brookhaven National Laboratory in June 1966, the nuclei data on U-238 for use in the Evaluated Nuclear Data File B (ENDF/B) are presented. The data cover the energy range from 0.001 eV to 15 MeV. Data sources are referenced, and the theoretical methods used in evaluating certain data are described. A complete listing of the data in the ENDF/B format is provided.

#6604066
Wong, C.Y.
EFFECTIVE MASS B, AND STIFFNESS PARAMETER C_F FOR QUADRUPOLE VIBRATION IN EVEN-EVEN SPHERICAL NUCLI
Nuclear Data, A:271-280 May 1968

Values of the effective mass $B$ and the stiffness parameter $C_F$ for quadrupole vibration of even-even spherical nuclei have been computed from the $B(E2)$ values and energies of first $2^+$ states adopted by Stekhop and Grodin. A comparison of these with values predicted by a liquid-drop model shows $B$ an order of magnitude larger than the predictions. The value of $C_F$ is usually less than the prediction but at magic numbers is many times greater.
Experimental and theoretical values of neutron spectra for a number of moderators and "poisons" in several geometries and at different temperature are presented graphically. The theoretical values are in point-kernel and moments-method shielding calculations. The cross sections were calculated from empirical expressions that permit the determination of fast neutron effective removal cross section for any element. Cross sections are presented for 83 elements.

In considering the possible physical and chemical effects of fission product elements at high temperatures on graphite and other materials, a table of data on the fission yields of the elements taken as the sum of the yields of all mass chains ending with a stable nuclide of the given element or a nuclide that has a half-life of one year or more was compiled. This gives a distribution of the elements, which corresponds approximately to that obtained from U-235 where a fuel element has run for an extended period of time (order of a year). The fission yields were taken from Schmitt and Zumwalt who have compiled a reasonably up-to-date summary of thermal neutron, U-235 fission yields. Also included in the table are the boiling points of the elements and qualitative data on the carbides. The volatility category of the elements is listed.
Atomic and Molecular Properties
The general transmissivity equation for computing slant-path molecular absorption spectra is developed and two methods for evaluating this equation, the direct integration and that which assumes a model of the band structure are discussed. Five band models are discussed and twelve methods for computing molecular absorption based on these band models are presented. Spectra computed by band-model methods are compared with spectra calculated by direct integration of the general transmissivity equation and with open-air field measurements of absorption spectra. Conclusions concerning the capability of band-model methods for predicting slant-path absorption spectra are stated and recommendations for future research are outlined. A summary of open-air field measurements and laboratory measurements of absorption spectra for homogeneous paths is presented, and a computer program for computing the equivalent sea-level path, the Curtis-Godson equivalent pressure, and the absorber concentration for atmospheric slant paths of any model atmosphere is given in appendix I.

#6900723
Anding, D.
BAND-MODEL METHODS FOR COMPUTING ATMOSPHERIC SLANT-PATH ABSORPTION SPECTRA
Report-No. 7142-21-T, Michigan University, Infrared and Optical Sensor Laboratory, Ann Arbor, Michigan February 1967
292p
Available from CFSTI, AD 815481

In this supplement, additional data are compiled on the optical materials, which were used for infrared instrumentation, treated in 2589-11-S, as well as data on three new materials, Irrran-1, Irrran-2, and diamond. In addition, several errors and ambiguities in the original report are corrected and clarified.

#690016
Ray, J. A.; Thompson, J. C.
ATOMIC AND MOLECULAR COLLISION CROSS SECTIONS OF INTEREST IN CONTROLLED THERMONUCLEAR RESEARCH
ORN-113 Revised, Oak Ridge National Laboratory, Oak Ridge, Tennessee January 1965
316p
Available from CFSTI

A graphical compilation is presented of atomic and molecular cross sections of interest to controlled thermonuclear research. The present compilation is a revision of ORNL-113, published in 1960. The cross sections are shown, as a function of energy, for collision processes involving molecular ion dissociation, charge exchange, excitation, ionization, photodissociation, scattered energy loss, secondary products and recombination. Pertinent nuclear cross sections are also included. A bibliography is given covering the literature since 1950.

#6811011, 6811012, 6811018, 6811021, 6811022, 6900215, 6902047, 6906024
Barnett, C. F.; et al.
135p, 202p, 257p, 176p, 225p, 264p, 288p, 245p
Available from CFSTI

These annotated bibliographies contain references of interest to atomic and molecular processes research. Bibliographical sources consist of more than eighty-five scientific journals and five abstract journals. The references are classified into categories with appropriate sub-categories. Each entry in the respective category is entered alphabetically by author and includes the reactants or the atomic or molecular system of interest.

#6500871
Bass, A. M.; Broida, H. P.
A SPECTROPHOTOMETRIC ATLAS OF THE SPECTRUM OF CH FROM 3000Ä TO 5000Ä
20p
Available from Superintendent of Documents, GPO

The near ultraviolet and visible emission spectrum of CH was recorded and presented in the form of a spectrophotometric atlas. The spectrum was recorded photoelectrically from an acetylene-oxygen flame in the region 4900 to 5000Ä by use of a high-resolution grating monochromator. Each of the lines in the CH spectrum is identified.
This compilation, which is designed primarily as a reference book for research scientists, is concerned with radiative and collisional processes involving atoms or molecules. It provides surveys covering the following topics: forbidden and allowed lines and bands, photoionization, photodetachment; recombination, attachment; elastic and inelastic scattering; recombination, energy levels in Rydberg states, excitation; recombination, energy levels by slow electrons; collision broadening; of spectral features; encounters between atomic systems including range, energy loss, ionization, attachment, radiative transfer, elastic scattering, mobility, diffusion, relaxation in gases, and chemical reactions. A chapter is devoted to the use of high temperature shock waves and accounts are given of the other main experimental methods. The relevant theoretical work is also described, detailed mathematics being avoided as far as possible.

This book is the result of a four-year survey by the X-ray Laboratory at the Johns Hopkins University. The primary objectives are to: (1) recompute all published wavelengths on a consistent basis; (2) include data published subsequently to the Sandstrom survey of 1957 and, in particular, refer to all wavelengths measured in the Johns Hopkins Laboratory; (3) convert these wavelengths to an absolute basis; (4) tabulate the data on which the recommended values are based; and (5) include explicitly probable errors for all adopted wavelength values. The $W$ K alpha, line peak was taken as a wavelength standard. The first part of the tabulated data includes wavelength, wavenumber, energy values arranged according to series and by element. The K, L, M, N and O series, as well as absorption edges are given. The second part of the tables presents wavelength and energy values in order of increasing wavelength. The appendix contains an evaluation of wavelength data and presents for each line a literature reference, a published, adjusted and recommended value, and reasons for making the adjustment, if any adjustment was made.

This publication contains two critical compilations: 1) X-Ray Wavelengths. Inconsistencies in accepted values (in $x$ units) of x-ray reference lines have recently been demonstrated, although all are supposedly based on "good" calcite crystals. Factors supporting the selection of the $W$ K alpha line as the X-Ray Wavelength Standard are critically discussed. A review is given of the experimental measurements which are used to establish the wavelength of this line on an absolute angstrom basis. Its value is $W$ K alpha $= 0.19091000 + 5$ ppm. The wavelengths of the Ag K alpha, Mo K alpha, Cu K alpha, and the Cr K alpha have been established as secondary standards with probable error of approximately one part per million. Six- or more x-ray lines have been used as reference values in a comprehensive review and reevaluation of more than 2700 emission and absorption wavelengths. The recommended wavelength values are listed in tables with probable errors; corresponding energies are given in keV. A second table lists the wavelengths in numerical order, and likewise includes their probable errors. A third table gives the recommended x-ray Atomic Energy Levels. All of the x-ray emission wavelengths have recently been reevaluated and placed on a consistent $W^*$ scale. For most elements these data give a highly overdetermined set of equations for energy level differences, which have been solved by least-squares adjustment for each case. This procedure makes "best" use of all x-ray wavelength data, and also permits calculation of the probable error for each energy difference. Photoelectric measurements of absolute energy levels are more precise than x-ray absorption edge data. These have been used to establish the absolute scale for eighty-one elements and, in many cases, to provide additional energy level difference data. The x-ray absorption wavelengths were used for eight elements and ionization measurements for two; the remaining five were interpolated by a Moseley diagram involving the output values of energy levels from adjacent elements. Probable errors are deduced from an absolute energy basis. In the original source of the present data, a table of energy levels in Rydberg units is given. Difference tables in units of Rydbergs, and milli-A* wavelengths, with the respective probable errors, are also included there.

This book contains the energy levels of atoms, and the differences between those levels, as computed from a table of x-ray wavelengths. The first set of tables contains values and probable errors of the energies of all occupied levels of each element, presented both in Rydbergs and in electron volts. The second set of tables contain differences between all the energy levels presented in the first set, in units of Rydbergs, electron volts, and milliangstroms.

This book is a reference for use in the analysis of organic and inorganic compounds through infrared spectra. The subject matter is confined to the cesium bromide region. The book is divided into four major parts: Part I discusses the characteristic frequencies and the conditions which must exist, the types of vibrations occurring in the infrared, some uses of the cesium bromide region, calibration standards, and sources of spectra in the cesium bromide region. Part II presents a complete review of the known characteristic frequencies in the cesium bromide region. Most of the common organic classes and some of the inorganic classes are covered. Part III contains the collection of 1566 infrared spectra covering the cesium bromide region. Part IV is a chemical class bibliography consisting of over 2000 references to literature on infrared data below 700/cm through 1966. The book also contains
The infrared absorption spectra of some 400 aliphatic and aromatic hydrocarbons have been investigated from 15 to 35 microns and the characteristic absorption frequencies incorporated into spectra-structure correlation charts. The classes of compounds studied were alkanes, alkenes, cyclopropanes, cyclobutanes, cyclopentanes, cyclohexanes, substituted benzenes, naphthalenes and biphenyls. The skeletal bending frequencies of the alkanes and alkenes and the nonplanar bending frequencies of cycloalkanes and aromatics are the most useful. The wavelength and intensity of the out-of-plane ring frequencies of aromatic molecules give some indication of the nature of the substituents. Typical infrared spectra of the hydrocarbons are presented.

The infrared absorption spectra of some 400 aliphatic and aromatic hydrocarbons have been investigated from 15 to 35 microns and the characteristic absorption frequencies incorporated into spectra-structure correlation charts. The classes of compounds studied were alkanes, alkenes, cyclopropanes, cyclobutanes, cyclopentanes, cyclohexanes, substituted benzenes, naphthalenes and biphenyls. The skeletal bending frequencies of the alkanes and alkenes and the nonplanar bending frequencies of cycloalkanes and aromatics are the most useful. The wavelength and intensity of the out-of-plane ring frequencies of aromatic molecules give some indication of the nature of the substituents. Typical infrared spectra of the hydrocarbons are presented.

A review of the work accomplished in the far infrared region is presented, and most of the available literature is cited. A double beam double pass spectrophotometer equipped with cesium bromide optics is described. This instrument extends the useful spectral range to 35 microns. The general features of the instrument's construction and its performance capabilities are discussed. Conventional infrared techniques were used, and in some instances simplified in obtaining infrared spectra. A final conclusion is derived. Many common solvents such as carbon disulfide, furan, benzene and dioxane, have large "open windows" in the long wavelength region, and the infrared spectra of these and other solvents are given. The advantages of far infrared spectra, and some of the special techniques of obtaining infrared spectra in the cesium bromide region, are also discussed.

Spectra of approximately 100 aromatic molecules, varying in size, shape, and structure, are assembled in the handbook. The list of molecules begins with the simple aromatic molecule benzene and includes larger and more complicated systems. Although a special effort has been made to include practically all of the currently popular organic scintillators, the choice of the other molecules has been rather arbitrary. The luminescence characteristics of each aromatic molecule have been measured in a systematic fashion. Fluorescence and absorption spectra are plotted for each molecule and additional data related to the fluorescence process are given in each graph. The additional data include the fluorescence decay time, the fluorescence quantum yield, the natural lifetime as computed from the absorption spectrum, the Stokes loss, the wavelength of the center of gravity of the fluorescence spectrum, and the average wavelength of the fluorescence spectrum. Such supplementary material as topical bibliographies, tables to convert wavelength to wave number and energy, a table of values of the index of refraction of compounds as a function of wave number, and a table of values of oscillator strengths have been placed in the Appendix.

This article surveys the restricted subject of properties of isolated atomic and small molecular negative ions. Included in the discussion are electron affinities, electronic states, optical properties, and the experimental and theoretical methods for studying these properties. Excluded from the discussion is the vast group of problems associated with large polyatomic negative ions and...
with most of the rich subject of collision processes involving negative ions, such as
detachment and transfer of electrons in collisions.

**#6500168**

*3.313*

Beynon, J. H.; Williams, A. E.

**MASS AND ABUNDANCE TABLES FOR USE IN MASS
SPECTROMETRY**

New York: Elsevier Publishing Company

1965

570p

These tables are intended to bring up to date those
previously published as an Appendix to "Mass Spectrometry and its Applications to Organic
Chemistry," by J. H. Beynon, 1960, and to extend
their range. They have been calculated and printed
on a Fortran Computer and are based on the
mass standard 12C = 12u exactly. The list has been
restricted to the four most common elements of
organic chemistry, carbon, hydrogen, nitrogen and
oxygen, and the combinations listed have been
restricted to containing at least one atom of
carbon and six or fewer atoms of nitrogen or
oxygen. The introduction to the tables is given in
English, German, French and Russian.

**#6500138**

*3.113*

Bhalla, C. P.

**TABLES OF ELECTRON RADIAL FUNCTIONS AND TANGENTS OF
PHASE SHIFTS FOR LIGHT NUCLEI** (2 = 1 THROUGH 10)

NBS Monograph 81, National Bureau of Standards,
Washington, D. C.

August 6, 1964

393p

Available from Superintendent of Documents, GPO

To facilitate the theoretical analyses of beta-
decay experiments in light nuclei, electronic
radial wave functions, evaluated at the nuclear
radius, and tangents of phase-shifts are tabulated
for total angular momentum equal to 1/2 and 3/2.
Separate tables for electrons and positrons are
given for ten values of 2, starting from 2 equal to
one, in steps of unity and for beta momentum values
from 0.1 mc to 42.0 mc in steps of 0.1 mc. The
nucleus is represented as a sphere with uniform
charge distribution. The nuclear radius, rho, is
taken to be 1.2A^1/2 x 10^-13 cm in the major body of
these tables. However, additional tables for 2 = 6
and 2 = 7 are given for rho = 1.1A^1/2 x 10^-13 cm
and rho = 1.3A^1/2 x 10^-13 cm.

**#6500824**

*2.113-3.113*

Bhanot, V. B.; Gupta, S. C.

**MASS-SPECTROSCOPIC ATOMIC MASS DIFFERENCES**

Nuovo Cimento, 10:1011-1023

June 1963

New mass-spectroscopic data on atomic mass
differences in the medium and heavy mass regions
were compiled and tabulated. Corresponding values in
the new scale based on C-12 as the standard are
also given. Unadjusted experimental values for
nucleic masses in the new scale were listed. For
a large number of stable nuclides in the region
from strontium to antimony the newer values
obtained from the data of Demikhov et al., are
more accurate than the corresponding values listed
in the recent exhaustive mass table of Koenig et
al. Attention is drawn to the advantages of the
isotopic doublets over the published conventional
mass doublets in the heavy mass region.

**#6711002**

*3.313*

Bieliski, B. H. J.; Gebicki, J. M.

**ATLAS OF ELECTRON SPIN RESONANCE SPECTRA**

New York: Academic Press

1967

665p

This atlas is made up of electron spin resonance
spectra recorded and published by several hundred
scientists during the last eighteen years. The aim
of the book is to make available those spectra of
interest to experimental spectroscopists, including
all spectra which could be assigned with reasonable
certainty to specific paramagnetic atoms, molecules
or ions. Spectra are made up of overlapping
signals due to two or more paramagnetic materials
unless the individual components could be
identified and the spectra assigned. A few
complicated spectra were omitted because the only
available pictures were not clear enough for the
resolution of all individual lines. All spectra of
oriented single crystals, which would require
much additional information, have been omitted.
Published results through late 1964 and some
unpublished spectra are included. The following
data are included with each spectrum: paramagnetic
species, irradiated material, formula for
paramagnetic species, method of preparation of
sample, medium, temperature, a list of lines observed,
relative line intensities, splitting constants, spin
density, and reference to original publications.

**#6601006**

*3.213*

Birnbaum, G.; Maryott, A. A.

**LINE SHAPES, WIDTHS, AND SHIFTS IN MICROWAVE
SPECTRA OF GASES**

Research Report No. 228, Hughes Research
Laboratories, Malibu, California

December 1961

34p

This report is a survey of results obtained in
recent years concerning the shape, width, and
frequency shift of microwave lines in dilute and
compressed gases. The dilute gas results are
concerned with relatively sharp lines associated
with molecular transitions and atomic hyperfine
transitions in hydrogen and optically pumped alkali
metal vapors in various buffer gases. The compressed
gas results are concerned with rather
broad lines or absorption bands and, in particular,
with the shape and width of the nonresonant spectra
of symmetric top molecules, frequency shifts in the
microwave spectrum of O2, NH3, and NB3, collision-
induced absorption in nonpolar molecules, and
absorption in the microwave wings of the infrared
rotational lines of HCl and DCl.

**#6908045**

*4.213-8.213-5.213*

Bondi, A. A.

**PHYSICAL PROPERTIES OF MOLECULAR CRYSTALS, LIQUIDS,
AND GLASSES**

New York: John Wiley & Sons, Inc.

1968

502p

This book develops a methodology enabling chemists
and chemical engineers to relate certain physical
properties of condensed phases to molecular
structure. The states of matter treated are
molecular crystals, liquids, and glasses. The
physical properties covered are P-V-T and related
thermal properties, and certain transport
properties. Energy of vaporization, van der Waals
dimensions, molar refractivity, dipole moment data
dielectric loss are used but not discussed
throughout. Table 4.1. page 414, the only table that
requires molecular structure increments. Background material presented is a guide to
the literature.

**#6805003**

*3.213*

Bovey, F. A.

**NMR DATA TABLES FOR ORGANIC COMPOUNDS, VOLUME I**

New York: Interscience Publishers

1967

610p

The purpose of this book is to provide a convenient
source of high resolution NMR data on a large
number and variety of organic compounds and a few
related inorganic compounds. These data can be
useful in two general ways:
1. to assist in the identification of new compounds and the recognition of known compounds by matching of spectral data;
2. to provide data on known compounds, in order to test new correlations, hypotheses and calculations.

This volume covers the literature from 1959 to 1969. The data for each compound are arranged in a horizontal series of entries which form columns for compound number, empirical formula, structural formula and name, solvent and concentration, temperature and reference compound, chemical shift and multiplicity, spin-spin coupling, spectrum type and/or analysis, observing frequency, and remarks and bibliographic reference.

#6805001 *3.013
Bramson, M. A.
Rodman, R. B. (Translator)
Wolfe, W. L. (Editor of Translation)
INFRARED RADIATION, A HANDBOOK FOR APPLICATIONS WITH A COLLECTION OF REFERENCE TABLES
New York: Plenum Press
1968
623p

This is the first volume of a series by M. A. Bramson which includes discussions on the basic principles of optics and the use of Planck's law for radiant intensity, for radiant existence and for radiant-transmission, terms of wavelength, frequency and photon number. The Wien and Rayleigh-Jeans expressions and modifications of the Stefan-Boltzmann law are derived. Chapters contain discussions on radiation, color, brightness temperatures, two-color emissivity-independent temperature-measurement methods, measurements of flames, bodies cooler than the radiometer, particle-transparent bodies, effects on measurements, and give tables of blackbody functions. This handbook includes procedures for computing the properties of infrared radiation over a temperature range from 100 to 6000°K. A collection of "Reference Tables for the Infrared Radiation from Heated Bodies" comprises the second portion of the book and includes tabulations of the Planck function, formulas derived from it, and the corresponding integrals. The book is aimed as a reference source and as a text for senior or graduate courses in infrared technology.

#6607004 *3.213
Brown, S. C.
BASIC DATA OF PLASMA PHYSICS
Cambridge, Massachusetts: The M.I.T. Press
1959
336p

This text consists of a combination of the class notes and the Basic Data technical report for a graduate course in gas discharges at M.I.T. put into book form. Its chapters deal with such topics as elastic-collision cross sections, charge-transfer cross sections, diffusion and mobility, inelastic collisions of electrons, diffusion-controlled breakdown, electron attachment, breakdown controlled by electron attachment, recombination, secondary electron emission by electrons, breakdown controlled by secondary electron emission, the second Townsend coefficient, D-C breakdown, the corona discharge, glow discharges, steady-state microwave discharges and arc discharges. The text has an author index as well as a subject index.

#6806002 *3.213
Bruegel, W. (Compiler)
NUCLEAR MAGNETIC RESONANCE SPECTRA AND CHEMICAL STRUCTURE, VOLUME 1: THE SPECTRAL NMR PARAMETERS OF COMPOUNDS WITH ANALYZED SPECTRA
New York: Academic Press
1967
235p

These tables contain the spectral nuclear magnetic resonance (NMR) parameters, i.e., the values of the chemical shift (magnetic shielding) and of the coupling constants of chemical compounds whose spectra have been analyzed. Section 1 of the tables deals with those cases where the atoms capable of resonance are either hydrogen or hydrogen together with other resonating nuclei, such as F-19 and P-31 (if they are present in a state where the spectrum can be described in a simple way, e.g., by reporting a single chemical shift of a nucleus other than hydrogen). All analyzed compounds of this type known to the reviewer are included with the exception of certain organic molecules containing Si, Ge and Sn, for which only some model-compounds are shown. Section 2 of the tables summarizes data on nuclei other than hydrogen. All inorganic compounds have been omitted.

#6908057 *3.113
Bucci, P.
PHOTONICS IN FAST NUCLEAR REACTORS, VI. APPENDIX
Ingeneersia Nuclleara, 8:33-77
November-December 1967

Tables, graphs and phase diagrams on the properties are presented. The characteristics of the systems Pu-Al, Pu-Be, Pu-Cr, Pu-Fe, Pu-H, Pu-Hg, Pu-Mo, Pu-Nb, Pu-O, Pu-Ta, Pu-Th, Pu-U, Pu-V, Pu-W and Pu-Zr are discussed and shown in graphic form.

#6512044 *3.313
Buckley, F.; Marvott, A. A.
TABLES OF DIELECTRIC DISPERSION DATA FOR PURE LIQUIDS AND DILUTE SOLUTIONS
NBS Circular 589, National Bureau of Standards, Washington, D. C.
November 1, 1958
Available from the Superintendent of Documents, GPO

Primary dielectric dispersion data and characteristic dispersion parameters are tabulated for almost 200 substances in the liquid state and for dilute aqueous and non-aqueous solutions with more than 150 solutes. There are 6 tables and 1 section of graphs. There are 4 tables for pure liquids, 2 containing summaries of the derived dispersion parameters and 2 containing the primary data. The section on graphs supplements the tables for pure liquids and contains reproductions of pertinent data that are available only in the form of graphs.

#6509014 *3.113
Buckmaster, H. A.
TABLES OF ANGULAR MOMENTUM TRANSFORMATION MATRIX ELEMENTS d_i,j (j = 2, 4, 6)
Canadian Journal of Physics, 42:386-391
February 1964

Crystalline electric field angular momentum operator equivalent transforms are given.

#6704004 *3.123
Candler, C.
ATOMIC SPECTRA AND THE VECTOR MODEL, SECOND EDITION
1964
412p

This second edition of Atomic Spectra compiles data and atomic spectral information through the use of the vector model. Where quantum mechanics has refined the predictions of the model, these predictions are compared with experimental results. MKS units are used entirely for all equations and data. The book contains chapters dealing with the hydrogen atom data, alkali doublets, the alkaline earth absorption spectra, the Zeeman effect, the Paschen-Back effect, the periodic system, the doublet laws, displaced terms, combination to several electrons, elements of the short periods,
long periods, the lanthanides, the actinides, line intensities, the sum rules and (jj) coupling, series limit, hyperfine structure andquadropole radiation. Tables of protonic, neutronic and deutonic nuclei, natural atomic units, the Grotrian diagrams, Rydberg term tables, and a bibliography are also included.

Carter, R. S.; Rush, J. J. (Editors) 
Molecular Dynamics and Structure of Solids 
NBS Special Publication 301, National Bureau of Standards, Washington, D. C. 
June 1969

Chapter 14: Molecular Dynamics and Structure of Solids

This volume is based on material presented at the Second Materials Research Symposium of the National Bureau of Standards, held October 16-19, 1967. It provides a review of the application of spectroscopic and diffraction techniques to the study of the structure and dynamics of molecular solids. Invited papers on the theory and practice of the major experimental methods, including neutron and x-ray diffraction, neutron inelastic scattering, infrared and Raman spectroscopy and nuclear magnetic-resonance, serve as a background for the technical presentation and discussion of results which follows. The topics covered in a series of invited and contributed papers include the lattice dynamics of molecular crystals, the spectroscopy and crystal structure of organic and inorganic solids and the dynamics of polymers.

Chapter 15: Applications of Spectroscopy

About 2800 articles of interest to mass spectroscopists are listed as a result of computer searching of 270,000 scientific articles published during 1966. The 149 papers presented at the 14th Annual Conference on Mass Spectrometry and Allied Topics, ASTM E-14, May 22-27, 1966, are also included. In addition to the bibliography, an author index and permuted title index (KWIC) are included.

Chapter 16: Special Topics

The present collection of 1000 infrared spectra was initiated with the intent of improving and adding to the sources that already exist in a variety of forms. The criteria set up to provide specifications for the evaluation of infrared spectra include three classes of spectra: critically defined physical data, research quality analytical spectra, and approved analytical spectra. The collection includes four sets of indices: alphabetical index, chemical classes index, molecular formula index, and a numerical index. The Chemical Abstracts System of nomenclature has been employed in naming the compounds and additional names have been added as cross-references to facilitate location.

Chapter 17: Summary and Conclusions

For purposes of evaluation, infrared spectra can be classified into three categories: Class I, standard spectra; Classes II and III, reference spectra. Class I spectra are designated as "standard spectra" because they are of a sufficiently high quality to be acceptable as physical constants of the substances under precisely defined conditions of measurement, and further refinements in spectrophotometric technique will not be expected to change them significantly. Few, if any, spectra of this quality have as yet been produced. Class II spectra are reference spectra obtained on the best currently available commercial infrared grating spectrophotometers, operated at maximum efficiency under conditions consistent with acceptable laboratory practice. The purity of the compounds must be rigidly specified with respect to the type and quantity of any spectrosopically detectable impurity present. Class III are the absorption spectra of individual compounds, and resins of known composition that have been obtained with sufficient accuracy to be useful in the identification of unknown materials. Class III is to be regarded primarily as a repository for usable data recorded prior to the general availability of high resolution grating spectrophotometers. The Board
of Management of the Cobalt Society recommends that spectra in Classes II and III be called "reference spectra" and that they be so designated in the published charts and in associated literature. It strongly recommends that the use of the word "standard" be reserved solely for spectra of Class I.

A two-volume table for converting wavelengths in standard air to wavenumbers in vacuum was computed by using the equation

$$\nu_{\text{vac}} = \frac{1}{(n^2 - 1)}$$

where n was computed from Edlen's 1953 equation for the refractive index of air. Wavenumbers are given to the nearest 0.001 cm\(^{-1}\) for wavelengths from 2000 to 7000 Å in volume I, and 7000 Å to 100 microns in volume II. Proportional tables are given for linear interpolation between entries of lambda. Also included are the vacuum increase in wavelength, (\(\eta - 1\)) and the refractivity of standard air in the form (\(\eta - 1\))\(x_{1000}\).

The propagation of electromagnetic waves through an ionized, gaseous medium can be described in terms of a set of macroscopic parameters that are a function of the physical properties of the medium, the permittivity, conductivity, and permeability. This report calculates the propagation parameters for the normal ionosphere above 50 km, and from these parameters calculates theoretical values for electron collision frequencies and scattering cross sections in the ionosphere.
Relative intensities of 39,000 spectral lines with wavelengths between 2000 and 9000 angstroms have been determined on a uniform energy scale for seventy elements. The light source was an arc between copper electrodes to which a single element was added in the ratio of one atom of element to 1000 atoms of copper. The temperature of the arc was determined, by comparison of the observed intensities with published relative gf-values, to be 5100 ± 110 K. The degree of ionization of eleven elements in the arc was determined by comparison of intensities in spectra of neutral and ionized atoms with known absolute gf-values. With the aid of Saha's ionization equation, the electron density in the arc was found to be 2.4x10^{17} cm^{-3}. The ionization of seventy elements separately added to the arc was then calculated with Saha's equation. A correction was made for diffusion of atoms from the arc stream. With this information, relative values of gf on a uniform scale can be computed for the 25,000 lines which have been classified. By calibration with known absolute gf-values, the results were put on an absolute scale. Absolute transition probabilities for 25,000 lines of 112 spectra have been calculated in this way and the results are tabulated by spectrum. The wavelengths in angstroms, energy levels to the nearest kajyer, g in 10^{-6} per second, gf, and log gf are given for each line.

A sixteen-inch double-focusing mass spectrometer employing the peak matching method of measurement has been used to measure the atomic masses of all stable isotopes in the region ruthenium to xenon. Atomic masses of 53 radioactive nuclei have been calculated from mass differences derived from nuclear reaction and beta-decay energies. Nucleon binding and pairing energies have been calculated from the resulting mass table. The effect of the shell closure at 2 = 50 on the systematics of nucleon binding and pairing energies has been investigated in greater detail than has previously been possible. The discontinuity in proton binding energy is shown to be caused by a decrease in binding energy of protons beyond 2 = 50. The main result of the study of nuclear systematics in this region seems to be that the binding energies of both neutrons and protons exhibit smooth behavior except for discontinuities at a shell closure. The presence of doubly charged, diatomic tellurium satellites which interfere with the singly charged ion peaks is also noted.
wide spectral region, from about 4370 angstroms to beyond the photographically accessible infrared. Fifty-four bands have been identified in the photographically accessible region. The tables list the wavelengths of the molecular lines in standard air, the intensities of the lines normalized to 100 for the strongest line in the band, the relative intensities on two discharge tube plates, the wave numbers of the lines in vacuum and the rotational quantum numbers of the various branches.

*Weiss, crystalline compounds of this.*

#6905034
Dickson, P. F.; Jones, M. C.
IMPURIFIED SPECULAR REFLECTANCES OF METALS AT LOW TEMPERATURES
Cryogenics, 8:24-29
February 1968

This paper graphically displays values, derived from the literature of the normal spectral reflectances ($\rho\nu$) of ten pure metals and an alloy at low temperatures. The metals and alloy included are aluminum, antimony, bismuth, copper, brasses, gold, silver, lead, tin, nickel and sodium.

*Dub, M. (Editor)*
New York: Springer-Verlag New York, Inc.
1966
856p

Volume I of the series Organometallic Compounds covers the derivatives of the transition metals of Group VIII of the Periodic Table. It is a comprehensive, noncritical source of information concerning organometallic compounds containing at least one carbon-metal bond. The information includes methods of preparation, properties, chemical reactions, applications, addition compounds and biological properties. This volume covers the literature from 1937 to 1964. In addition to the tables of data and information given, a bibliography of over 2300 references and a listing of review publications and monographs are included.

*Dub, M.; Weiss, R. W. (Editors)*
ORGANOMETALLIC COMPOUNDS: METHODS OF SYNTHESIS, PHYSICAL CONSTANTS AND CHEMICAL REACTIONS, VOLUME II: COMPOUNDS OF GERMANIUM, TIN AND LEAD INCLUDING BIOLOGICAL ACTIVITY AND COMMERCIAL APPLICATION, SECOND EDITION, COVERING THE LITERATURE FROM 1937 TO 1964
New York: Springer-Verlag New York, Inc.
1967
727p

Volume II of the series Organometallic Compounds covers organic derivatives of germanium, tin and lead. It is a comprehensive, noncritical source of information concerning organometallic compounds. The scope is limited to compounds containing at least one carbon-metal bond. The information includes methods of preparation, properties, chemical reactions, applications, addition compounds and biological properties. This volume covers the literature from 1937 to 1964. In addition to the tables of data and information given, a bibliography of over 2300 references and a listing of review publications and monographs are included.

#6903046
Dukelskii, V. M. (Editor)
COBBIE, D. M. (Translator)
ELECTRONIC AND ATOMIC COLLISIONS BIBLIOGRAPHY OF RUSSIAN LITERATURE FOR THE YEARS 1946-1966
Atomic and Molecular Processes Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee October 1968
69p

This bibliography is based on a machine translation of a Russian edition. The original Russian references and their translations are both listed. The references are grouped into major categorical processes. The numerical order is unchanged from the original Russian bibliography. The major categories include general collisions, electron-atomic collisions, collisions of atomic particles and their applications. An author index is provided.

#6906009
Eisenberg, D.; Kaumann, W.
THE STRUCTURE AND PROPERTIES OF WATER
New York and Oxford: Oxford University Press
1969
296p

In this book the authors summarize the many experimental observations on water and present theoretical discussions which effectively correlate the data. The topics covered include the water molecule and forces between water molecules; the thermodynamic properties of steam; the structures of the classical and nonclassical water; the thermodynamic, electrical, spectroscopic, and transport properties of the ices and of liquid water; hydrogen bonding in ice and water; and models for liquid water. The authors emphasize the relation of the properties of ice and water to their structures, and discuss properties such as infrared and Raman spectra because they reveal a great deal about these structures. The inclusion of some background material in physical chemistry makes the book accessible to the biologist, biochemist, and geologist as well as the chemist and physicist. An addendum listing a number of recent articles on the structures and properties of water completes the volume.

#6508065
Feor, G.; Gorter, C.-J.; Smits, L.-J.
TABLES OF CONSTANTS AND NUMERICAL DATA, VOLUME 7, SELECTED CONSTANTS, DIAMAGNETISM AND PARAMAGNETISM, PARAMAGNETIC RELAXATION
Paris: Masson & cie, Editeurs
1957
318p

In French, introduction in French and English

These tables list constants and numerical data for materials which exhibit diamagnetism or paramagnetism. These materials include the elements, binary alloys, mineral compounds, double or complex salts, organic compounds, free radicals, hydrides, oxides, hydracids, acids of mineral oxides, organic acids, anhydrous single and double salts, hydrates, anhydrous single and double salts, complex anions in order of decreasing valency, complex anions in order of increasing valency, non-ionic and metallo-organic compounds. The properties given for these materials are magnetic moments, Curie points and Curie constants, mean atomic susceptibility, moments calculated according to Weiss's law, magnetic anisotropy of crystalline compounds, molecular susceptibilities, effective moments, atomic moments, and other related data. The study of paramagnetic relaxation information both on the interactions between paramagnetic ions and on the interactions between the ions and the crystalline lattice. The data give magnetic techniques required for the production of very low temperatures by adiabatic demagnetization of paramagnetic substances. Cross references and bibliographical references are included.
This is a compilation of ionization and appearance potentials of positive ions published from 1955 through June 1966. The compilation lists the ion formed, the parent species from which it was formed, the other products of the process, the threshold energy for the formation of this ion, and the method by which this data was obtained. Where feasible, the heat of formation at 298 K of the positive ion has been computed for each entry using auxiliary thermochemical data. From these computed values "best" values have been chosen.

Available from Superintendent of Documents, GPO

---

Nuclear Science Abstracts was searched from Volume 7 (1955) through Volume 12 Number 15 (August 15, 1966) for material on the elastic scattering of alpha particles on alpha particles. The references are arranged numerically within each volume with the abstracts from the most recent volumes presented first.

Available from the Superintendent of Documents, GPO

---

A tabulation of attenuation coefficients of X-rays and gamma-rays from 0.01 to 100 MeV for 29 materials is presented. A summary of information on the probability of the basic interaction processes of photons with matter and a detailed analysis of experimental and theoretical evidence are included. Present information on the basic processes is adequate for many applications; however, improved theory and additional experimental data are needed in certain areas. A comparison of calculated and experimental coefficients points up this need.

Available from Superintendent of Documents, GPO

---

This bibliography contains lists of selected references on preparation methods, crystal structure, transport properties, and physical and mechanical properties for the Groups IV, V, and VI transition metals. An abstract is included along with each reference entry.

Available from Superintendent of Documents, GPO
This document is a bibliography on ion-molecule reactions of chemical interest. The bibliography is listed in a chronologic-alphabetic arrangement and includes 431 titles and references taken from the published literature since 1900. The coverage is limited to reactions between ions and molecules leading to changes in the atomic composition of either the projectile ion or target molecule. No attempt has been made to completely cover resonant charge exchange or elastic scattering.

This monograph deals with the principles and applications of chemical analysis by flame photometry, and data obtainable by useful for flame analysis. The topics dealt with in this section are: basis of flame analysis, apparatus and methods of measurement, control of the flame and introduction of the sample, optics and equipment, the flame emissions, standardization and measurement and flame analysis errors. The applications dealt with include agricultural applications, and biological applications, with applications to geology, industry and other fields. The data in the appendix include the intensities of wavelengths of 3400 lines and bands of 145 atoms, ions and molecules representing 70 elements that can be determined by flame photometry. This information was taken from results of spectrographic analyses published in the literature. The elements are arranged by emitting species in one table, and by wavelength in the other.

Experimental results pertaining to atomic and molecular charge-transfer processes reported in the literature from 1950 through 1960 have been collected and tabulated on IBM punched cards. The parameters listed for each reaction are: the energy interval of the measurement, the maximum charge-transfer cross section, and the energy at which this maximum is reached. In such cases as the Massey criterion (\(\Delta G/\Delta H\) approximates 1 for maximum \(\Delta G\)) still represents the best method for correlating existing data and for making at least rough predictions for other charge-transfer events, calculated values of \(\Delta G\) have been included. The report also contains a fairly complete bibliography of the charge-transfer literature, including both experimental and theoretical papers. The punched cards from which the tables in this report were printed are available for duplication.

This monograph provides a five-year supplement to the original Infrared Absorption Spectra Index for 1958-1962. The literature coverage of this Index includes 66 journals and one collection of spectra in book form. It contains approximately 70,000 references to Infrared absorption spectra published during 1958-1962. As in the previous volume of this series, each reference indicates the journal, volume, and page where an actual spectrum of the compound is reproduced.

This monograph provides in the form of an index for locating nuclear magnetic resonance (NMR) and electron spin resonance (ESR) spectroscopy data in the literature. It includes more than 8000 references to NMR and ESR spectra published in six volumes worldwide journals and collections of spectra during 1958-1963. Each reference in this index indicates the journal, volume, and page where an actual spectrum of the compound being sought is found. The index is arranged alphabetically according to the compounds whose spectra are given.
This textbook covers the field of diatomic molecule spectra, and contains tabulations of the electronic levels and the vibrational and rotational constants of all known diatomic molecules, plus photographs of certain important molecular spectra. The topics covered in the text portion are: a review of the elements of atomic spectra, observed molecular spectra, rotation and vibration of diatomic molecules, discussion of electronic states and transitions, details of electronic states and transitions, building up principles, electron configurations, continuous and diffuse molecular spectra, examples, results, and applications. The data tabulated include the electron term, vibrational constants, rotational constants, dissociation energy, internuclear distance, and observed transitions in every known state of every diatomic molecule.

This text is the second volume in a series of three on Molecular Spectra and Molecular Structure and deals with the information obtained from infrared and Raman spectra. A large number of illustrations and tables have been included in which theoretical results are summarized or observed data are collected. The tables contain critically reviewed data published through 1945. The chapters in this text deal with rotation and rotational spectra, vibrations, vibrational energy levels, vibrational eigenfunctions, vibrational infrared and Raman spectra, interactions of rotation and vibration, rotation-vibration spectra and applications. The book contains an appendix of physical constants and conversion factors, a bibliography of almost a thousand references, and an author and subject index.

This book presents a comprehensive review of theory and experimental results of studies dealing with electronic spectra of polyatomic molecules. The text deals with the following subjects: (1) electronic states, electronic energy, vibrational and rotational structure of states, effects of electron and magnetic fields on energy levels; (2) electronic transitions: types of transitions, rotational and vibrational transitions, Zeeman and Stark effect; (3) building-up of principles: correlation of electronic states, electron configurations, stability of molecular electronic states; (4) dissociation, predissociation and recombination, continuous and diffuse spectra; (5) electronic spectra of individual molecules and their interpretation for molecules of three to twelve atoms. The data presented include energy-level diagrams and individual molecular spectra as well as tabulated data of molecular constants of most polyatomic molecules (up to 12 atoms) for which discrete spectra have been analyzed. Constants tabulated for each molecule are: point group, electronic levels, vibrational frequencies, rotational constants, structural parameters, observed transitions, and literature references.

Ultrasound and visible absorptions of organic compounds are characteristic of absorbing systems that consist of a so-called chromophore group and the auxochrome groups bonded to it. For this reason, the data in this volume are organized from the point of view of the absorbing system. Data contained in Table I include: absorbing system, compound name, solvent, wavelength of maximum absorption, molar absorptivity, and reference to the literature. Table II is a list of absorbing systems arranged by wavelength of maximum absorption which can be used for identifying compounds by absorption maxima.
Hubbell, J. H., Berger, M. J.

PHOTON ATTENUATION
Engineering Compendium on Radiation Shielding, Vol. I, Chapter 4, pages 167-202
Berlin: Springer-Verlag 1968

"Photon Attenuation" is a section of the Engineering Compendium on Radiation Shielding which tabulates Photon cross sections and attenuation coefficients. The Photon Attenuation results are for 83 element and 18 element tabulations at 10 keV and 10 MeV. The Photon Attenuation table is based on a new analysis of all available information. Discussions of the Photon cross sections underlying the interaction coefficients are included.

Hughes, R. L.; Smith, I. C.; Lawless, E. A.; Holzmann, R. T. (Editor)

PRODUCTION OF THE BORANES AND RELATED RESEARCH
533p

This is a review and analysis of borane chemistry. The topics covered include: precursors to the boranes; formation of the boron-hydrogen bond and diborane; preparation and properties of the boranes, alkylboranes, and carbaboranes; kinetics and mechanism studies; preparation and properties of pentaborane and decaborane from diborane; separation and purification; analytical methods; boron-nitrogen chemistry acid-base chemistry; and appendices on toxicity; infrared spectra, molecular structures, physical properties of boranes and their derivatives.

This report updates and extends previous NBS tabulations. Section 1 contains the information of most immediate practical use: (a) a tabulation of the attenuation coefficient for 25 elements (U-1 to U-92) between 10 keV and 100 GeV and (b) a tabulation of the energy absorption coefficient for 50 elements, and 18 elements between 10 keV and 10 MeV, and for air, water, and 7 elements up to 100 MeV. Section 2 contains detailed information on the predominant processes (photoelectric absorption, Compton scattering, and pair production) and a brief discussion of other processes which combine to give the attenuation coefficient. Theoretical and experimental data are reviewed, and auxiliary tables and approximate formulas are given. Section 3 contains tabulations of cross sections for the predominant processes between 10 keV and 100 GeV for 23 elements and for 13 compounds and mixtures.

Hubbell, J. H. et al.

PHOTON CROSS SECTIONS, ATTENUATION COEFFICIENTS, AND ENERGY ABSORPTION COEFFICIENTS FROM 10 keV TO 100 GeV
NSRDS-NBS-29, National Bureau of Standards, Washington, D. C.
August 1969 83p

Available from the Superintendent of Documents, GPO

This is a directory of international scientists engaged in or interested in the field of electronic, atomic, and molecular collisions; it is a revision of the 1965 directory and is a publication of the Atomic and Molecular Processes Information Center.

International Union of Pure and Applied Chemistry, Commission on Molecular Structure and Spectroscopy

INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY, PROCEEDINGS OF THE SECOND INTERNATIONAL CONFERENCE ON NUCLIDIC MASSES, VIENNA, AUSTRIA, JULY 15-19, 1963
New York: Springer-Verlag 1964 480p

Separate abstracts were prepared for 30 of the 37 papers included. Six other papers were previously abstracted for NSA 17:31400, 32998, 33950, 36639; 18:1057; and 19:14607. The remaining paper describes the interest of the International Union of Pure and Applied Chemistry in the values of the fundamental constants.

Jones, R. N.; et al.

COMPUTER PROGRAMS FOR ABSORPTION SPECTROPHOTOMETRY
NRC Bulletin No. 11, National Research Council of Canada, Ottawa, Canada 1968 159p

Nine "building block" computer programs for performing the basic numerical computations of absorption spectrophotometry are described and listed. They are written in FORTRAN IV for card input and output. The input and output formats are standardized to permit easy interfacing to produce more complex data processing systems. Though developed for infrared spectrophotometry these programs are adaptable for use in the visible and ultraviolet. The operations covered include scale change and scale interval change, location of band maxima and minima, smoothing and differentiation, slit function convolution and deconvolution, single band generation, profile analysis and asymmetry analysis of single bands from the truncated moments, and ordinate addition.
This report contains all the published atomic emission lines contained in the literature up to January 1967. Each line is divided into two main sections: the emission lines by element and a finding list. The entries give the element and a wavelength, intensity, classification, and the references for the line. This report is divided into four sections, each dealing with the same elements. Section I is a list of intense lines by ion; Section II is a finding list of these intense lines; Section III is a list of all the lines arranged by element; Section IV is a finding list for all elements. Sections I and II are those lines of Section III which are reported as having high intensities. Section III is the most complete list, containing 30,000 lines. Section IV has the weaker lines of Section III removed and contains 23,000 lines. The lines marked with an asterisk in Sections III and IV are those listed in Sections I and II. Sections I and III include the data on ionization potentials (I.P.) given in Charlotte Moore's Atomic Energy Levels, (National Bureau of Standards Circular No. 467). The intensities given in the tables are those published for spark sources. Throughout this table, the original author's estimates for intensity are used.

This bibliography was compiled by the JILA Datal Center for the purpose of providing a list of published reports which contain information on measurements of electron collision cross sections. The data for molecules are listed in arbitrary order.

This bibliography provides references to comprehensive collections of low energy atomic collision data. The data sought are those for wave-length regions of continuous absorption, absorption and ionization coefficients, from which cross sections can be derived, as well as references to molecular absorption in wave-length regions where there is continuous absorption even though there are line absorptions superimposed. This report contains a section which describes the data in the references of the
This compilation of critically evaluated data presents basically two kinds of data. The first is "gross" of "total" ionization cross-sections, which correspond to the probability of producing a single positive charge from any atomic or molecular species regardless of the mass or multiplicity of change on the ions produced. The other kind is that which gives the relative probability of producing ions with a specific charge and mass (in the case of molecules where dissociation can take place). The tables of "total" cross-section data are grouped by their Z-number. The data for seven molnuclear species are ordered by their atomic weights. After these tables, the absolute cross-section data for single ionization by electron impact are presented. There then follows a section of cross-sections for the dissociative ionization of diatomic molecules, and finally a section giving relative numbers of multiple-charged ions. Accuracy of data is in all cases at best 5%.

A comprehensive compilation of low energy electron collision cross-section data is presented. This is the first of three parts and is limited to experimental measurements, including data for all atomic species and for those molecules which are important in aeronomy, astrophysics and plasma physics. Literature data through September of 1968 are included. The compilation is divided into three main groups: ionization, including all processes which result in electron ejection from atomic or molecular targets; dissociation, including processes which lead to ionized fragments; and vibrational and rotational excitation. The data are organized by species in the order: atoms, diatomic molecules, triatomic molecules, etc. The atoms are ordered by the Z number, and the molecules are ordered according to the atom with the largest Z in the molecule.

This is the second part of a comprehensive compilation of low energy electron collision cross-section data. The compilation is limited to experimental measurements and includes data for all atomic species and those molecules which are important for aeronomy, astrophysics, and plasma physics. The data included were taken from literature published through December 1968. However, reference to some materials is obtained from abstracting journals, so this compilation probably does not include all data published in the late fall of 1968. All of the cross section data in this volume of the compilation are for electronic excitation. The data are organized by initial species in the following order: atoms, diatomic molecules, triatomic molecules, etc. The atoms are ordered by the Z number of the nucleus; the smallest Z's first. For molecules, the number of the atom with the largest Z in the molecule is taken to characterize the molecule. The molecules are then ordered by these characteristic Z's, smallest Z's first. If necessary the second highest Z atom in the molecule is used in the same way to determine the order. If the line emitted is characteristic of the ionized species, the data are located following the neutral species.
This index contains data on 3200 mass spectra. The data tabulated for each spectrum are the compound name, molecular weight, peak mass/charge ratios and the intensities of these peaks. The spectra were taken from the American Petroleum Institute Project 44 files, the ASTM files, and the Manufacturing Chemists Association Project. Each entry contains the serial number by which the spectra may be found in the files. Compounds are arranged by molecular weight and by the mass/charge ratio of the four highest peaks.

This list includes 16,130 compounds, the ultraviolet or visible spectra of which have been indexed by the Wyndotte - ASTM Punched-Card Index as issued through the fifth supplement. The serial numbers given provide a reference to the location of the original spectra as published in the literature or in standard catalogues of spectra. Arrangement is by the order of the numbers of atoms in the molecular formulas. A journal and book reference appendix is included. CODEN has been used for journal references.

This section of the Landolt-Bornstein tables contains data on the fundamental physical constants and the properties of atoms and ions. Information for the use of the tables, including data on systems of measurement and units, has also been included. A discussion of atomic and molecular spectra, including wavelength standards, electron terms, important spectral line wavelengths, ionization potentials, electron affinities, X-ray spectra, Zeeman and Stark effect data, pressure broadening and pressure displacement of spectral lines, and oscillator strengths and lifetimes of excited states is presented. Other properties of atoms and ions compiled include electron distribution, X-ray scattering and absorption, cross sections, magnetic moments, magnetic and electric polarizability, and Faraday effect data.

This section of the Landolt-Bornstein tables contains atomic physical data of molecules which characterize the structure and dynamics of their nuclear framework. The following properties have been tabulated: atomic distances and structures,
valence energies of chemical bonds, dissociation energies of chemical bonds, and molecular vibrations and rotations.

#6500527 *0.1151-2.2051-3.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME I: ATOMIC AND MOLECULAR PHYSICS, PART 3: MOLECULES II (ELECTRON SHELL) AND AN APPENDIX TO VOLUMES I, PARTS 1, 2, AND 3 Eucken, A.; Hellwege, K.-H. (Editors) Berlin, New York: Springer-Verlag 1951 772 p

In German

This section of the Landolt-Bornstein tables contains tables on the electron shell of molecules. The volume includes data published through September 1951. The following properties have been tabulated: band spectra of diatomic molecules, electron band spectra of multatomic molecules, light absorption of solutions in the infrared and visible region, ionization energies of molecules, optical rotary power of molecules, electrical and optical polarizability of molecules, magnetic moments of molecules, diamagnetic polarizability of molecules, and quantum output of photochemical reactions.


In German

This section contains data on the atomic physical interpretation of crystal physics as of November 1954. The following properties have been tabulated: the characteristics of lattice types, structures, crystal dimensions of inorganic and organic crystals, atomic and ionic radii, lattice energies, integral vibrations of crystals, electron emission, energy bonds, X-ray spectra and bonding states, electron spectra of crystals, and high frequency spectra of crystals.

#6500529 *0.1151-2.2051-3.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME I: ATOMIC AND MOLECULAR PHYSICS, PART 5: ATOMIC NUCLEI AND ELEMENTARY PARTICLES Hellwege, K.-H. (Editor) Berlin, New York: Springer-Verlag 1952 470 p

In German

This section of the Landolt-Bornstein tables contains tables on atomic nuclei and elementary particles. These tables were compiled by the authors from literature on these subjects published through November 1951. The properties tabulated are: the hyperfine structure of atomic terms and atomic lines, a table of all atomic nuclei and their properties, mass spectrometric measurements of nuclear reaction energies, electron line spectra of artificially radioactive atoms, the radioactive families, nuclear level schemes of radioactive transformations, energy schemes of the light atomic nuclei, beta-spectra, naturally radioactive atoms and their alpha- and beta-spectra, scattering of charged particles by atomic nuclei, nuclear transformation processes, passage of alpha-particles, electrons, protons and gamma-rays through matter and cosmic radiation.


In German

This section of the Landolt-Bornstein tables contains tables and figures for the optical properties of substances. Data are compiled from literature published through June 1962. The data are compiled for: the optical constants and reflection of metals, alloys, and thin metal layers; refractive indices of inorganic and organic materials as a function of wavelengths; optical constants of selected solid materials in broad wavelength regions; rotation of the polarization plane in crystals; double refraction; piezoelectrical and electrooptical constants of piezoelectric crystals; light refraction by glasses and transparent plastic; light permeability, double refractive index; water; refractive index and optical rotatory power of pure liquids and solutions; Cotton-Mouton effect of liquids; double refraction due to Kerr effect in liquids; Kerr constants of liquids; refractive indices of gaseous elements and compounds; refraction and dispersion of air and of hydrogen compounds; Cotton-Mouton effect of gases and Kerr constants for gases.

#6500537 *0.1151-2.2051-3.151 LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME III: ASTRONOMY AND GEOPHYSICS Bartels, J.; Bruggencate, P. T. (Editors) Berlin, New York: Springer-Verlag 1952 795 p

In German

This volume of the Landolt-Bornstein series contains data on astronomy and geophysics. The material has been taken from literature published through 1951. The astronomical section includes data on astronomical instruments, position and time determination, abundance of the elements in the universe, the mechanics and physics of the planets, moons, comets and meteors, stellar radiation, stellar spectra, stellar positions and movements, double stars, variable stars, novae, the Milky Way, star clusters and extragalactic nebulae. The section on geophysics contains data on the earth's gravity and tidal forces, radioactivity density; elastic, electrical, magnetic and thermal properties of minerals and rocks; seismicity and earthquake waves; properties of the earth's interior, surface and magnetism; physical and chemical properties of sea water; sea tiles, hydrography, meteorology and climatology; and upper atmosphere physics.


In German

This section of the Landolt-Bornstein tables includes data on metallographic principles and
This section of the Landolt-Bornstein tables deals with the technological aspects of electrical, light, and X-rays. It is compiled from the literature published through 1956. Wherever possible, the compilers have drawn from several values available for a given quantity, and the values they feel is the most accurate. The properties tabulated in the electrical section are electrical conductivity, resistance and temperature coefficient of resistance; conductivity of molten metals; effect of pressure, tension, and cold machining on electrical resistance; resistance as functions of frequencies; resistance of thin layer semiconductors; thermoelectric emf of metal pairs and thermodiffusion factors; effects of pressure and cold treatment on thermodiffusion; conduction of electricity in gases, breakdown potentials of gases; insulating liquids, filling and impregnating media, fibrous material insulators, insulating foils, mica insulators, rubber insulators; resin, plastics, quartz, glass, ceramic materials and rocks as insulators; and magnetic materials. The quantities tabulated in the light section are: sensitivity of the human eye, illuminating materials, light sources, light filters and photographic films. The X-ray section includes data on measurements and units of X-ray technology.

The various types of angular correlation coefficients have been tabulated in the literature. These tables provide the coefficients for a complete calculation of the basic angular correlations in alpha-, beta-, and gamma-spectroscopy. In chapter 2, the angular correlation formalism is given in a necessarily incomplete collection of relations involving the tabulated coefficients, i.e., 3j-symbols, 6j-symbols, 9j-symbols, F-coefficients, and GAMMA-coefficients. In the computer tables these symbols are given in upright capital letters: $J = j, J_3 = j_3$, etc. and $\Gamma = \Gamma_j$. These quantities are given solely by angular momentum conservation for the total system in consideration. When answering the question of how the angular momenta of the various subsystems of particles may combine to fulfill the conservation condition, one ends up with Clebsch-Gordan-coefficients for angular momenta and with Racah-coefficients for more than two. To achieve a higher degree of symmetry, Wigner has rewritten both types of coefficients as the 3j- and 6j-symbols, respectively. In more complicated correlations (e.g. triple correlations, where the intermediate transition also is observed) a new combination of 9j-symbols appears which is usually called the F-coefficient. The GAMMA-coefficient is a slightly modified form of this F-coefficient useful for beta-spectroscopy. The coefficients mentioned above are extensively tabulated in this volume. For the 9j-symbol, however, only selected ones are listed. In addition, there is a chapter dealing with angle functions. Since angular
correlations are generally developed in a series of Legendre polynomials, these functions should be very useful.

#6605007
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW SERIES
GROUP II: ATOMIC AND MOLECULAR PHYSICS, VOLUME 1: MAGNETIC PROPERTIES OF FREE RADICALS
Fischer, H.; Hellwege, K.-H.; Hellwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1965
154p
In German and English

This book contains data on the magnetic properties of free radicals, including inorganic and organic uncharged free radicals and radical ions. These data were taken from published results of ESR and atomic beam resonance experiments. The data compiled include: name and structural formula of a free radical, magnetic susceptibility of a crystal matrix or solvent used for the radical, temperature of measurement, frequency of measurement, g-factor, hyperfine splitting parameter, primary references for the values quoted, and further references on the same subject.

#6706020
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW SERIES
GROUP II: ATOMIC AND MOLECULAR PHYSICS, VOLUME 2: MAGNETIC PROPERTIES OF COORDINATION AND ORGANOMETALLIC TRANSITION METAL COMPOUNDS
Koenig, H.; Hellwege, K.-H.; Hellwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1966
578p
In German and English

Magnetic properties are listed for coordination and organo-metallic transition metal compounds. The magnetic susceptibility section lists for each compound an average susceptibility and magnetic susceptibilities at a particular temperature, magnetic moments and paramagnetic Curie temperatures, and the experimental method used to obtain data. The magnetic resonance section lists for each compound the spectroscopic splitting g-factor, fine structure and hyperfine structure parameters, half-width of the EPR line, and temperature and frequency of measurement.

#6706036
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW SERIES
GROUP II: ATOMIC AND MOLECULAR PHYSICS, VOLUME 3: LUMINESCENCE OF ORGANIC SUBSTANCES
Schnillen, A.; Legler, R.; Hellwege, K.-H.; Hellwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1967
416p
In German and English

The most important data on the luminescence of over 1000 luminescent organic substances investigated have been assembled into a main table. Most substances are shown not only in a pure state (liquid or solid) but diluted in several solvents as well. On compiling the literature, it was noticed that not all data were of equal reliability, e.g. owing to insufficient purity of the substances investigated or to the application of theories which had become out of date in the meantime. For these reasons the main table only contains the physical magnitudes which are known with sufficient reliability for the majority of the substances investigated. Data appearing in the main table for each compound include: its state, solvent used, if any, temperature, wavelength used for stimulation, absorption and emission wavenumbers for fluorescent and phosphorescent emission, and data on infrared and other chemical properties.

This collection provides data on the rotation of molecules in the gaseous state obtained by microwave spectroscopy. The following molecular constants are presented: rotational constants, centrifugal distortion constants, rotation-translation interaction constants, beta-type doubling constants, and isotopic masses or mass ratios. These constants are separately tabulated for diatomic molecules, linear molecules, symmetric top molecules, and asymmetric top molecules. Also presented are electric dipole moments, nuclear quadrupole moments, and hindered rotation constants. There are also diagrams of the spectroscopic splitting of the electric dipole moment, for those molecules whose structure could be deduced from the microwave measurements with sufficient reliability. References are tabulated with a special table for magnetic constants determined by Zeeman effect.

#6802001
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW SERIES
GROUP II: ATOMIC AND MOLECULAR PHYSICS, VOLUME 5: MOLECULAR ACOUSTICS
Schaffs, W.; Hellwege, K.-H.; Hellwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1969
286p
In German and English

This volume lists data on acoustics based essentially on the molecular properties of matter, and values of sound velocity, absorption coefficients, and dispersion data. The measurements were made at frequencies ranging from 10 kc to 200 Mc. Preference was generally given in compiling data to measurements for which statements on density, chemical purity, and the temperature coefficient were added. Often, however, several statements can be found for a measured value. Generally, the limit of accuracy is 1% to 2%. Data are given on sound velocity in gases and liquids; for pure gases and liquids as a function of temperature, pressure, and structure; also for mixtures of gases, mixtures of liquids, and solution. Data are compiled on sound absorption and dispersion in gases; sound absorption in homogeneous liquids, in solution, and on sound propagation in polymers, also on sound propagation in isotropic or quasi-isotropic solids (metals, inorganic noncrystalline solids). There are special sections on sound propagation in liquid helium and shock wave velocity.
This volume of the Landolt-Bornstein New Series contains data useful in the fields of astronomy and astrophysics. The following topics are covered: optical instruments, radio-astronomical devices, performance of telescopes, photoelectric photometry, position and time determination, astronomical constants, abundances of the elements in the universe; physical and mechanical properties of the sun, planets, satellites, comets, meteors, artificial earth satellites, interplanetary space, stellar positions and motions, stellar spectra, star colors and magnitudes, physics of stellar atmospheres, stellar structure and evolution, double stars, variable stars, novae and supernovae, planetary nebulae, star clusters and associations, properties of the nearest stars, galactic structure, kinematic and dynamical properties of the galaxy, interstellar space, galaxies, and clusters of galaxies.

This book introduces a continuing series of collections of absorption spectra in the ultraviolet and visible region for a wide range of compounds, all constructed to the logarithmic scale. Observed log I/I values have been entered into tables containing the original measured data. Spectrum characteristics unobtainable with the necessary degree of accuracy from the graph are thereby made accessible. Although the work was started in Hungary, subsequent volumes contain spectra by authors of several countries. This introductory book presents the fundamental theoretical information required for the interpretation of the spectra, as well as conversion tables: wavelengths - wave numbers, wave numbers - wavelengths, wave lengths - frequencies, wavelengths - cm/mole, transmittancy - absorbancy - absorbptancy, list of the formulas used, and spectroscopic conversion table for energies. The book includes discussions on structure of spectra, application of the Franck-Condon principle to polyatomic molecules, anharmonicity, dissociation and predissociation, the form of electronic band systems, chromophores, complexes, the role of the medium effect, and a bibliography.

These volumes contain a description of the absorption spectra between 2000 and 5000 A of recently synthesized compounds, of compounds known in the literature, of data observed for individual compounds in different solvents, of spectra compiled for the investigation of pH relationships of spectra of isomer pairs and of a few vapor spectra. Absorption spectra are given for 2065 compounds. The logarithm of the molar extinction coefficient is plotted as a function of wavelength. A tabulation of optical densities is also given for each compound as a function of wavelength. Volume 1 is the first edition, Volumes 2 and 3 are the second edition, and Volumes 4 through 11 are the first edition.

This book contains series of theoretically calculated multicomponent symmetrical ESR spectra. These spectra are for systems of 4 to 9 components having binomial or certain intermediate intensities, for both lorentzian and gaussian line shapes of the components.

This compilation is a collection of results of gas chromatographic data taken from the literature on the subject. The tables can be used, either for identification of unknown materials, or for selecting a column for separating a known mixture. Data have been tabulated on the basis of the liquid phase used and the serial number of the reference material.

This book of nuclear quadrupole coupling constants is divided into two sections. In the first, the basic theory of quadrupole coupling is presented, the methods available for the experimental measurement of the coupling constants reviewed and the relationship between them and a molecule's electronic structure discussed both from the rigorous and approximate viewpoint. In the second, the experimental data are reviewed and compiled by means of the approximate methods developed in the first section. This includes the halogen coupling constants of organo-halogen
compounds, N quadrupole coupling constants, constants of compounds of the typical elements of the Periodic Table, compounds of the transition and rare-earth elements, and charge transfer complexes. Author and subject indexes are included.

McClellan, A. L.
TABLES OF EXPERIMENTAL DIPOLE MOMENTS
San Francisco: W. H. Freeman and Company 1963
71sp
This book is a non-critical compendium of experimental dipole moments taken from literature published from the 1920's through 1961. These dipole moments were derived principally from measurements of the dielectric constants of materials or microwave measurements. The materials are divided into three classes; compounds without carbon, compounds containing carbon, compounds of unspecified formula. The second class is the largest by far and contains about 6000 entries. For each compound, the formula, compound name, and one or more measured values for the dipole moment are given. For each measurement, the state of the sample and/or type of solvent used, temperature of measurement, method of measurement, and a reference number referring to the book's bibliography are given. The appendix contains a list of average dipole moments of 75 common chemical compounds, the available moments for twelve homologous allene series, and the criteria and treatment of data applied during the computation of the recommended values.

McGinnies, R. T.
X-RAY ATTENUATION COEFFICIENTS FROM 10 KEV TO 100 MeV
10p
Available from the Superintendent of Documents, GPO
A revision is given of the X-ray attenuation coefficients presented in National Bureau of Standards Circular 583. Table 4 of that publication is eliminated, and a new table is given for each material for photon energies less than 100 keV. The uncertainties in the estimates of attenuation coefficients at low energies are from 3 to 5 percent, which is the same as was previously given for higher energies. The cross sections for scattering are exchanged. Two values are listed for the photoelectric cross section, one calculated from the Sauter-Stobbe formula and the other derived from new experimental evidence. The procedures for smoothing experimental data are described and are generally the same as were used in Circular 583. In addition to the systematic coverage of the region from 10 kev to 100 Mev, some data are included for a number of elements based on experimental measurements below 10 kev and above 100 MeV. A comparison is made between calculated and experimental total attenuation coefficients at energies above 10 MeV.

McLafferty, F. W.
MASS SPECTRAL CORRELATIONS
Advances in Chemistry Series No. 40, American Chemical Society, Washington, D. C. 1963
117p
This table provides the empirical and structural formulas of ions that might be found at a particular mass/charge ratio in a mass spectrum, plus an indication of how each such fragment ion might have arisen. An indication of the probability of each such classification of ions occurring is also given. Thus, the table indicates possible ion structures and precursor molecules for each of the prominent ions in the mass spectrum of an unknown compound, with a further indication of the general probability of their occurrence. Over 4000 spectra are listed.

Mclntosh, W. H.; et al.
CORRELATION OF X-RAY CROSS SECTIONS SECTION II
UCRL-50174, California University, Lawrence Radiation Laboratory, Livermore, California January 1967
96p
Preliminary graphs and tables of X-ray cross sections in the range of 1 kev to 1 Mev are given. Theoretical data for magnetic moments have been recalculated using the Hartree-Fock equation and the fit to derived experimental photoelectric cross sections. Form factors available in the literature were used to compute the scattering cross sections. Where solutions to the Hartree-Fock equation were unavailable, Newilou's values for the Thomas-Fermi model were used. L and H shell jump ratios were calculated from the theoretical values of Rakavy and Ron. The graphs for each element show the experimental data points used to obtain the fit.

Mallmann, R. de; Suhner, F.
TABLES OF CONSTANTS AND NUMERICAL DATA, VOLUME 3: MAGNETIC ROTATORY POWER (FARADAY EFFECT) KERR MAGNETO-OPTIC EFFECT
Paris-V: Hermann & Cie, Depositaire 1951
138p
In French
This set of tables gives data on gases and vapors of simple compounds, mineral compounds, and organic compounds; on liquids of simple compounds, mineral compounds, and organic substances including hydrocarbons, esters of inorganic acids, alcohols, phenols, ethers, aldehydes, ketones sugars, acids, chloride acids, anides, anhydrides, nitrates, nitrates, azoimide, hydrazines, quaternary ammonium salts, heterocyclic compounds, and organo-metallic compounds; on electrolytic solutions of acids, bases, and mineral salts; and on molecular rotations of pure substances and electrolytic solutions. The Kerr magneto-optic effect section includes tables of rotation of ferromagnetic substances, graphs of rotation versus wavelength, dispersion of the rotation of visible light, dispersion of rotation of infrared, ellipticity, and rotation of non-ferromagnetic metals. The book contains formula indexes as well as bibliographical references.

Maryott, A. A.; Buckley, F.
TABLE OF DIELECTRIC CONSTANTS AND ELECTRIC DIPOLE MOMENTS OF SUBSTANCES IN THE GASEOUS STATE
20p
Available from the Superintendent of Documents, GPO
Values of the dipole moments and other pertinent information are tabulated for approximately 350 substances in the vapor state. All values derived from measurements of dielectric constants have been recalculated by one of two systematic procedures in order to place the work of various investigators in a more comparable basis than exists in the literature. Values obtained independently from microwave spectroscopy and other methods are also included. Values of the dielectric constants recommended for use are listed for helium, hydrogen, oxygen, argon, air, nitrogen and carbon dioxide. These selected values were derived from a consideration of radio frequency, microwave, and optical data. Much of the data given in this
The "static" dielectric constants of more than 800 substances in the liquid state were critically examined and tabulated in concise form. The table consists of three sections: A, Standard Liquids; B, Inorganic Liquids; and C, Organic Liquids. An indication of the probable accuracy of the data is given. Wherever feasible, a simple analytical function is employed to express the variation of dielectric constant with temperature.

This volume compiles optical rotatory power data from results published in the literature for steroids, listed according to their molecular formulae. The data given include the chemical formula, structure, specific rotatory power, name and concentration of solvent, temperature at which the measurement took place, and wavelengths. An alphabetical index of substances, author index, and list of bibliographical references are included.

The table have been superseded by #6809016 (NBSRDS-NBS-10).

Table of Dielectric Constants of Pure Liquids

Maryott, A. A.; Smith, E. R.

Available from the Superintendent of Documents, GPO

This volume compiles optical rotatory power data from results published in the literature for steroids, listed according to their molecular formulae. The data given include the chemical formula, structure, specific rotatory power, name and concentration of solvent, temperature at which the measurement took place, and wavelengths. An alphabetical index of substances, author index, and list of bibliographical references are included.

#651204, 6512005

Navrodineau, R.

Bibliography on Analytical Flame Spectroscopy, Volumes 1 and 2

Elicitott City, Maryland: Central Library of Spectroscopy

1965, 1966

64p, 54p

The collection of references on analytical flame spectrometry presented in these two volumes cover the period 1800 to 1966. Subject indexes, keyed by number to the references cited, precede most of the sections.

The tables of constants and numerical data in the literature for triterpenoids, listed according to their molecular formulae. The data given include the chemical formula, structure, specific rotatory power, name and concentration of solvent, temperature at which the measurement took place, and wavelengths. An alphabetical index of substances, author index, and list of bibliographical references are included.

Mathieu, J. P.; Ouirisson, G.

Available from the Superintendent of Documents, GPO

This volume compiles optical rotatory power data from results published in the literature for triterpenoids, listed according to their molecular formulae.

#6508064

Mathieu, J. P.; Ourisson, G.

Available from the Superintendent of Documents, GPO

This volume compiles optical rotatory power data from results published in the literature for steroids, listed according to their molecular formulae. The data given include the chemical formula, structure, specific rotatory power, name and concentration of solvent, temperature at which the measurement took place, and wavelengths. An alphabetical index of substances, author index, and list of bibliographical references are included.

Mathieu, J. P.; Petit, A.

Available from the Superintendent of Documents, GPO

This volume compiles optical rotatory power data from results published in the literature for steroids, listed according to their molecular formulae. The data given include the chemical formula, structure, specific rotatory power, name and concentration of solvent, temperature at which the measurement took place, and wavelengths. An alphabetical index of substances, author index, and list of bibliographical references are included.

#6508062
Wise, properties and i compil function in construction thorough SPECTRAL GENERAL a various the number the 10 wavelength 474, spectra C. methods compact increasing of the the high reactions effects, part calculating F. is front large revised supplementary JANUARY Miles, bibliography. The A uniform of other Standards, are adjusted and arranged TABLES Meggers, bibliographies. The Monograph 32, Part I and Part II, National Bureau of Standards, Washington, D. C. December 29, 1961; October 2, 1961. The relative intensities or radiant powers of 39,000 spectral lines with wavelengths between 2000 and 9000 angstroms have been determined on a uniform energy scale for seventy chemical elements. This was done by mixing 0.1 atomic percent of each element in powdered copper, pressing the powder-mixture to form solid electrodes which were burned in a 10 ampere, 220 volt direct-current arc, and photographing the spectra with a stigmatic concave grating with a step sector rotating in front of the slit. The sectored spectrographs facilitated the estimation of intensities of all elements relative to copper which were then calibrated on the energy scale provided by standardized lamps, and all estimated line intensities were finally adjusted to fit this calibration. Comparisons with other intensity measurements in individual spectra indicate that the National Bureau of Standards' spectral-line intensities may have average errors of 20 percent but first of all, they provide uniform quantitative values for the seventy chemical elements commonly determined by spectrochemists. These data are presented by element in part I, and all 39,000 observed lines are given in order of wavelength in part II.

Wavrodineau, R.; Boiteux, H. FLAME SPECTROSCOPY New York: John Wiley & Sons, Inc. 1965 721p. This text discusses analytical spectroscopy by flame. Topics emphasized range from the principles important to the proper selection, design, and construction of excitation sources, to the identification, interpretation, and understanding of the various radiations excited by flames, and to the relevance of certain reactions involved in the radiation processes. A large volume of data from the literature has been included with proper references. The book is divided into three parts. The first part consists of an analysis of various properties of flames, and of the description of the elements involved in the construction of the excitation source and of its function conditions. The second part explains the mechanism of spectra of the chemical species excited in the acetylene flames, and discusses some of the reactions occurring in flames. The third part comprises reproductions of spectra from various chemical species excited in the acetylene flames, followed by wavelength tables and two complementary bibliographies.


R-6272, Rocketdyne, Canoga Park, California August 12, 1965 222p. This report describes theoretical studies of the basic radiative properties of high temperature gases. The primary emphasis of this work has been devoted to methods of calculating spectral absorption coefficients of electronic transitions in diatomic molecules. The smeared line model has been used to calculate absorption coefficients in presented on a critically selected basis. To keep the bibliography to a compact size, a number of comprehensive papers which contain numerical results throughout extended iso electronic sequences are listed in a separate section. Also included in this bibliography is a supplementary list of selected important papers dealing with the subject of transition probabilities from a general point of view.

Milek, J. T. FLUOROCARBON GASES, DATA SHEETS DS-142, Hughes Aircraft Company, Electronic Properties Information Center, Culver City, California November 1964 111p. A compilation of the electrical properties of various halocarbon or halogenated hydrocarbons known as Freons, Genetrons, Arcons, etc., is presented. A master identification chart relating the trade names and numbers to the chemical name is included for easy reference. Detailed electrical properties include: properties, corona effects, dielectric constant, dielectric strength and dissipation factor. Each property is compiled over the widest possible range of pressure, temperature, electrode geometry, effects, and electrodes from references obtained in a thorough literature search. Physical and chemical property data are also included as electrical and electronic applications.

Miller, J. T. SULFUR HEXAFLORIDE DATA SHEETS DS-140, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California October 1964 68p. A compilation of the electrical properties of sulfur hexafluoride, a dielectric gas, is presented. Electrical properties include: properties, corona effects, dielectric constant, dissipation factor and dielectric strength. The latter property data section is segregated into parameter effects as follows: properties, corona effects, dielectric constant, dissipation factor and dielectric strength. Each property is compiled over the widest possible range of parameters obtained in a thorough search of the world's literature.


Miller, R. V.; Golden, S. A. RADIATIVE PROPERTIES OF GASES, VOLUME 1. GENERAL DISCUSSIONS R-6272, Rocketdyne, Canoga Park, California August 12, 1965 222p. This report describes theoretical studies of the basic radiative properties of high temperature gases. The primary emphasis of this work has been devoted to methods of calculating spectral absorption coefficients of electronic transitions in diatomic molecules. The smeared line model has been used to calculate absorption coefficients in presented on a critically selected basis. To keep the bibliography to a compact size, a number of comprehensive papers which contain numerical results throughout extended iso electronic sequences are listed in a separate section. Also included in this bibliography is a supplementary list of selected important papers dealing with the subject of transition probabilities from a general point of view.


#6500429 *3.153 Miller, J. T. FLUOROCARBON GASES, DATA SHEETS DS-142, Hughes Aircraft Company, Electronic Properties Information Center, Culver City, California November 1964 111p. A compilation of the electrical properties of various halocarbon or halogenated hydrocarbons known as Freons, Genetrons, Arcons, etc., is presented. A master identification chart relating the trade names and numbers to the chemical name is included for easy reference. Detailed electrical properties include: properties, corona effects, dielectric constant, dielectric strength and dissipation factor. Each property is compiled over the widest possible range of pressure, temperature, electrode geometry, effects, and electrodes from references obtained in a thorough literature search. Physical and chemical property data are also included as electrical and electronic applications.

#6500431 *3.153-4.153 Milek, J. T. SULFUR HEXAFLORIDE DATA SHEETS DS-140, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California October 1964 68p. A compilation of the electrical properties of sulfur hexafluoride, a dielectric gas, is presented. Electrical properties include: properties, corona effects, dielectric constant, dissipation factor and dielectric strength. The latter property data section is segregated into parameter effects as follows: properties, corona effects, dielectric constant, dissipation factor and dielectric strength. Each property is compiled over the widest possible range of parameters obtained in a thorough search of the world's literature.


Miller, R. V.; Golden, S. A. RADIATIVE PROPERTIES OF GASES, VOLUME 1. GENERAL DISCUSSIONS R-6272, Rocketdyne, Canoga Park, California August 12, 1965 222p. This report describes theoretical studies of the basic radiative properties of high temperature gases. The primary emphasis of this work has been devoted to methods of calculating spectral absorption coefficients of electronic transitions in diatomic molecules. The smeared line model has been used to calculate absorption coefficients in presented on a critically selected basis. To keep the bibliography to a compact size, a number of comprehensive papers which contain numerical results throughout extended iso electronic sequences are listed in a separate section. Also included in this bibliography is a supplementary list of selected important papers dealing with the subject of transition probabilities from a general point of view.
fifty such transitions. Ancillary work has included studies of methods of calculating energies of potential functions of the Rydberg-Klein-Rees type, solutions to the Schrödinger equation with such potential functions, and computational procedures for evaluating Franck-Condon and r-centroid factors.

#6500146
Mohler, F. L.
PHOTOIONIZATION OF ATOMS AND MOLECULES
NBS Technical Note 131, National Bureau of Standards, Washington, D. C.
January 1962
43p
Available from CFSTI, PB 161 632
This is a review of experimental results on photoionization of atoms and some molecules. There are some quantitative data on all the alkaline earths, magnesium, calcium and thallium and all rare gases except xenon. Results are given for the common gases: hydrogen, nitrogen, oxygen, C0, C02, NO, N2O, H2O and CH4. Autoionization - excitation to a state above the ionization threshold followed by transition to the ionized state - can be an important factor. Often the broad autoionization lines mask the true continuum. There are some mass spectrometry measurements of photoionization products for most of these molecules.

#6812003
*5.153-3.153
Moiseiwitsch, B. L.; Smith, S. J.
ELECTRON IMPACT EXCITATION OF ATOMS
August 1963
120p
Available from Superintendent of Documents, GPO
The experimental and theoretical literature about the electron impact excitation of atoms is reviewed. Theoretical methods ranging from the Bethe and Bloch approach to the distorted wave approximations are discussed and compared. Where possible, on theoretical grounds or through intercomparison, the reliability of the methods is discussed. A general critique of the optical method of measuring excitation functions is given, with the objective of promoting higher quality future experimental work. A critical study of existing experimental work is included. The conclusion is that most workers have ignored important physical and instrumental effects, and it may be presumed that the data in the literature are subject to many unrecognized systematic errors. The literature on the alkaline earths, rare gases, mercury, cadmium and zinc is surveyed but the quality of the literature does not support critical review. Beyond this some general comments about the physics of these atoms.

#6500118, 6500119
*5.153
6500120
Moore, C. E.
ATOMIC ENERGY LEVELS, VOLUME I. III TO 23V,
VOLUME II. 41N TO 72H
VOLUME III. 42M TO 57 La, 72Hf TO 89Ac
NBS Circular 467, National Bureau of Standards, Washington, D. C.
June 15, 1940; August 15, 1952; May 1, 1958
309p, 272p, 245p
Available from Superintendent of Documents, GPO
This series of three volumes is a critical compilation of atomic energy levels prepared at the National Bureau of Standards from the analyses of optical spectra. Volume I contains data on the spectra of hydrogen, deuterium, tritium, helium, lithium, beryllium, boron, carbon, nitrogen, oxygen, nitrogen, neon, sodium, magnesium, aluminum, silicon, phosphorus, sulfur, chlorine, argon, potassium, calcium, strontium, titanium, and vanadium (III to 23V). Volume II analyzes the spectra of chromium, manganese, iron, cobalt, nickel, copper, zinc, gallium, germanium, arsenic, selenium, bromine, krypton, rubidium, strontium, yttrium, zirconium, and niobium (24Cr to 41Nb). Volume III includes the spectra of molybdenum, technetium, ruthenium, rhodium, palladium, silver, cadmium, indium, tin, antimony, tellurium, iodine, xenon, cesium, barium, lanthanum, hafnium, tantalum, tungsten, rhenium, osmium, iridium, platinum, gold, mercury, thallium, lead, bismuth, polonium, radon, radium, and actinium (42M to 57La, 72Hf to 89Ac).

#6500023
Moore, C. E.
THE ATOMIC SPECTRA OF THE RARE EARTHS: THEIR PRESENCE IN THE SUN
JULY 1963
A summary of the present state of analysis of the first and second spectra of the rare-earth elements is presented with an extensive bibliography. The astrophysical importance of these spectra is stressed. Counts are given of the number of lines of each rare-earth spectrum identified at present in the accessible solar spectrum. Most of them appear only in the singly ionized state; exceptions are Eu, Tm, Yb, and Y. The possible presence of the strongest laboratory line of the rare-earth atomic spectra in the solar spectrum is suggested. If correct, this is the first evidence of a third spectrum line in the accessible range of the solar spectrum (λ > 3000 Å).

#6502038, 6905015
6906025, 6910015
Moore, C. E.
BIBLIOGRAPHY ON THE ANALYSIS OF OPTICAL ATOMIC SPECTRA,
SECTION 1: III TO 23V,
SECTION 2: 41N TO 72H
SECTION 3: 42M TO 57La AND 72Hf TO 89Ac
SECTION 4: 57La TO 72Lu AND 89Ac TO 99Es
September 1968; February, May, August 1969
90p, 63p, 43p, 55p
Available from the Superintendent of Documents, GPO
The three published volumes on "Atomic Energy Levels", NBS Circular 467, contain for each element of the periodic table a list of atomic energy levels, including the wavelengths, wave numbers, and quantum numbers of the levels and line transitions. The bibliography is being published in four sections; each covers the same elements as the respective volumes of AEL. Section I contains reference lists for the elements I through 23, corresponding to Volume I. The spectra of a given element are listed in order of increasing stage of ionization. Listings are included for spectra of the 23 elements. The original papers have been excised for nearly all of the quoted references. A selected list of general literature references is also included. Section II is similarly arranged, giving references to the spectra of the elements 24A through 41B, corresponding to AEL Volume II. For a given element the spectra are listed in order of increasing stage of ionization. The original papers have been examined for nearly all of the quoted references. Section III is similarly arranged giving references to the spectra of the elements, 42M through 57La and 72Hf through 89Ac, similar to AEL Volume III. For a given element the spectra are listed in order of increasing stage of ionization. The original papers have been examined for nearly all of the quoted references. Section IV concludes this publication, and completes the coverage of the Periodic Table. The references listed have been selected on a general basis as those needed for the preparation of Volume IV of
"Atomic Energy Levels". Since rare-earth structure is revealed in the new portion, 89Ac, the references for these elements are repeated in the present Section, which contains the two groups of elements: $^7$La to $^{71}$Lu and $^{89}$Ac to $^{232}$Th. A selected list of actual literature references is also included. Nearly all of the quoted references have been checked from the original papers.

#6500125

Moore, C. E.
MULTIPEL TABLE OF ASTROPHYSICAL INTEREST
PART 1: TABLE OF MULTIPELTS;
PART 2: FINDING LIST OF ALL LINES IN THE TABLE OF MULTIPELTS
NBS Technical Note 36, National Bureau of Standards, Washington, D. C.
November 1959
140p
Available from CFSTI, PB 151395

Pending the completion of a current edition, the 1945 Multiplet Table is being reprinted here to meet continuing demands. The leading lines in 196 atomic spectra of 55 chemical elements are listed in related groups called multiplets. Estimated intensities, excitation potentials and multiplet designations are given for the individual lines, and the multiplet as a whole. An extensive bibliography covers the source material used for the compilation. The Table is presented in two parts: Part I includes the multiplets, with the spectra of each element being given in order of increasing ionization, and the elements in order of increasing atomic number. Part II is a Finding List in which all the lines in Part I are entered in order of increasing wavelength, with their multiplet numbers. The range of the Table is from 2091A to 13164A. A supplementary table of "Forbidden Lines" extends from 2072A to 12645A.

#6500161

Moore, C. E.
SELECTED TABLES OF ATOMIC SPECTRA, ATOMIC ENERGY LEVELS AND MULTIPELTS, SI II, SI III, SI IV
NSRDS-NBS-3, Section 1, National Bureau of Standards, Washington, D. C.
June 25, 1965
37p
Available from the Superintendent of Documents, GPO

The compilation contains tables of atomic energy levels and multiplet tables for the spectra of Si II, Si III, and Si IV. The data in these tables were taken from literature articles published through 1965. Section A, Atomic Energy Levels, contains tables of energy levels of each atom or ion. The data given for each level include its electron configuration, spectroscopic notation, the value of the quantum number J, the energy expressed in cm$^{-1}$, and the interval in cm$^{-1}$ between levels of the same configuration and different values of J. The Multiplet Table, Section B, contains the wavelength, intensity and multiplet number of each multiplet found. Also listed are the excitation potentials, spectroscopic configurations, and values of J for the two levels between which the transition takes place. Literature references are given for both tables.

#6600020

Moore, C. E.
SELECTED TABLES OF ATOMIC SPECTRA, ATOMIC ENERGY LEVELS, AND MULTIPELTS, SI
NSRDS-NBS-3, Section 2, National Bureau of Standards, Washington, D. C.
November 30, 1967
22p
Available from the Superintendent of Documents, GPO

The present publication is the second section of a series being prepared in response to the increasing demand for a current revision of two sets of tables containing data on atomic spectra as derived from analyses of optical spectra. Both the atomic energy levels and the multiplet table are included in the same publication, as parts A and B, respectively. The sections are being prepared at irregular intervals for these spectra whose analysis are especially complete. A flexible printing system permits the arrangement of the various sections by atomic number, regardless of the order in which the spectra are published in the present series. This publication includes three spectra of silicon, $Z = 14$: Si II, Si III, Si IV. The present section contains similar data for Si I. The form of presentation is described in the text to Section 1, and need not be repeated here.

#6611020

Moore, C. E.; Minnaert, H.; Houtgast, J.
The SOLAR SPECTRUM 2035A TO 8770A, SECOND REVISION OF ROWLAND'S PRELIMINARY TABLE OF SOLAR SPECTRUM WAVELENGTHS
NBS Monograph 61, National Bureau of Standards, Washington, D. C.
December 1966
340p
Available from the Superintendent of Documents, GPO

The present compilation of solar spectrum wavelengths and intensities is essentially a second revision of Rowland's Table, corrected and supplemented by comparison with the photometric Atlas of the Weigert-Bode Hilger Photometric Atlas. Approximately 24,000 lines are listed. In a number of cases new wavelengths were determined. Measured equivalent widths from the Atlas were corrected for asymmetry and estimated line intensities recorded in the 1928 revision. From these directly measured equivalent widths have been derived reduced widths, which, if necessary, were corrected for dispersion in the solar spectrum. The intensity behavior of atomic lines in the solar spectrum is indicated by letters denoting strengthening, weakening, and trivial. Additional lines present only in the spot spectrum are also included, 237 in all. Revised identifications of the lines, as to chemical origin, are given for both atomic and molecular lines. For classified atomic lines the lower excitation potential and multiplet number are listed. For molecular lines the rotation branch and quantum number, and the vibration band are indicated. Note numbers refer to notes in which the complete designation of the band is given. An introductory text gives a detailed description of each column of the solar ledger. Figures are included to illustrate the procedure used to derive the observed equivalent widths $\Delta$ (A) and the reduced widths $\Delta$ (F). Tables include counts of lines of each spectrum recorded in the identification column, leading lines in the first row of each spectrum, and subgroups of molecules and elements present in the sun. About 73 percent of the lines are wholly or partially identified. Sixty-three elements are recorded at present. A number need further study. The number of molecules identified in the sun totals 11.

#6500121-6500124

Moore, C. E.
AN ULTRAVIOLET MULTIPEL TABLE, SECTION 1: H I TO 25V; SECTION 2: 240+ TO 41NH; SECTION 3: 420+ TO 750A; SECTION 4: FINDING LIST FOR SPECTRA OF THE ELEMENTS HYDROGEN TO NIOBIUM (Z = 1 TO 41); SECTION 5: FINDING LIST FOR SPECTRA OF THE ELEMENTS MOLYBDENUM TO LANTHANUM (Z = 42 TO 57) AND HAFNIUM TO BISMUTH (Z = 72 TO 88)
NBS Circular 488, Sections 1 to 4
Available from the Superintendent of Documents, GPO

An Ultraviolet Multiplet Table, NBS Circular 488, Table 1, a table of multiplet spectral lines, with wavelengths of 300A or less. These tables were taken from results of spectral observations, published in the literature. Sections 1 and 2 (6500121) contain spectra of neutral and ionized
atoms of Z = 1 to 41. Section 3 contains spectra of atoms of Z = 42 to 57 and 71 to 80. These tables are arranged, first by atomic number then by ionization series, and finally by wavelength. The data included in these tables are the wavelength and intensity of the multiplet line, the excitation potentials for the upper and lower states of the transition involved, the J-values and multiplet designation of the lines, and a literature reference for each line. Sections 5 and 6 give lists for the multiplet tables, arranged in order of wavelengths. Section 4 covers Z = 1 to 41 and Section 5 covers Z = 42 to 57 and Z = 71 to 80. These lists give for each line its wavelength, the radiating atom and ion, and the multiplet number.

#6809007 *3.153

65p Available from the Superintendent of Documents, GPO

This publication is a reprint of Appendix A of the book by Paul W. Merrill entitled, "Lines of the Chemical Elements in Astronomical Spectra" Carnegie Inst. Washington, Publ. 610, (1956). It contains partial Grottrian Diagrams of selected spectra of astrophysical interest, which give wavelengths, multiplet numbers and key letters for the transitions shown. The diagrams are accompanied by tabular keys in which the key letters indicate related lines in spectra similar in structure to those illustrated in the diagrams. There are 39 diagrams of spectra between hydrogen and nickel and tabular keys for 90 spectra between lithium and rhenium.

#6604027 *3.153

The radiative corrections to the high-energy Bremsstrahlung and pair-production spectra are calculated in the Weissacker-Williams approximation. Results for the soft-photon radiative correction to the spectra are given, and for the case of pair production the soft-plus-hard-photon radiative correction is also obtained. The radiative correction to the total pair-production cross section is found to be practically independent of the photon energy and of the atomic number of the target material. Comparison with available experimental data shows essential agreement within the relatively wide experimental limits.

#6902008 *4.153-3.153-3.153

This is a text on the properties of boron and boron compounds containing ten chapters, each written by experts in the different phases of boron chemistry. The chapters discuss the element boron, compounds of high boron content, borates, boric acid, boron reference. Sections 4 and 5 are listing lists for the multiplet tables, arranged in order of wavelengths. Section 4 covers Z = 1 to 41 and Section 5 covers Z = 42 to 57 and Z = 71 to 80. These lists give for each line its wavelength, the radiating atom and ion, and the multiplet number.


The Mossbauer Effect Data Index is an organized index to experimental research results relating to the Mossbauer Effect published through 1965. The index groups experiments first by nuclear transition (daughter isotope and gamma-ray energy), then by source material or host, and then by absorber (or scatter) material or host. Each experiment is described and summarized on an 80-column IBM card. The data listed for each experiment are: Mossbauer isotope and gamma-ray transition energy, host material in which the source atoms are imbedded, source temperature, absorber, absorber temperature, line width, dip of the spectrum (percentage change in counting rate from on-resonance to off-resonance), shift of the absorption spectrum, quadrupole splitting energy (for transitions between spin 3/2 and 1/2 levels), shape of the absorption spectrum and a code for the bibliography reference. A summary sheet is presented for each isotope presenting simplified decay scheme gamma-ray and x-ray energies, half-life and total internal conversion coefficient of the Mossbauer transition, natural isotopic abundance, and magnetic and quadrupole moments. There is an index to references by topic and author.

#6500449 *2.153-3.153

Available from Superintendent of Documents, GPO

The Compton energy versus angle relationship and the differential and integral Klein-Nishina cross sections are presented graphically as functions of the energy and direction of the scattered photon and of the recoil electron. These graphs are intended to serve the purpose of tables. Unpolarized primary gamma rays in an energy range from 10 kev to 500 Mev are considered. The accuracy of all curves is estimated at 1 percent. The advantage of this form of presentation is the convenience and accuracy of two-way interpolation. In general, interpolated values may be obtained with an accuracy of 2 percent.

#6809016 *3.153-3.153

Available from the Superintendent of Documents, GPO

This table revises, brings up to date, and extends the coverage on numerical values of dipole moments which was included in NBS Circular 537, Tables of Dipole Moments in the Gas Phase, 1964. Extensive comments are given on the definition of dipole moment and principal methods of dipole moment measurement, as well as an exposition of the criteria employed in selecting the tabulated data.
This paper reviews some aspects of a continuing research program in which intensity measurements have been made on a number of important and common diatomic molecular band systems, including the calculation of Franck-Condon factors, the \( \gamma \)-centroids, and the integrated emission intensity.

The book consists of a set of tables presenting reduced matrix elements and related quantities for all possible configurations of equivalent \( p, d, \) and \( f \) electrons. Particular attention is given to the recalculations of results already available in the literature, but most of the results on \( f \) electrons are new. The tables contain listings of states, fractional parentage coefficients, electrostatic energy matrices, and unit tensor operator matrices.

Schematic representations are given for both the formation of positive ions and the formation of negative ions in ionized air. The formation of positive ions can be divided into three regions. Region I contains the ionization of neutral constituents and two-body reactions in dry air. Region II contains reactions involving ionic species which are formed by three-body reactions. Region III contains reactions involving water vapors and shows the formation of the hydrated hydronium ions. The formation of negative ions proceeds predominantly through 01. The clustering of neutral elements to \( O \) gives additional channels for the formation of \( NO \), an ion which is not attacked by atomic oxygen. Rate constants are given, including the experimental values on which they are based, for the reactions in the schematic representations.

A table of the wavelengths of the rare-earth elements and several associated elements has been compiled for the use of spectrosco"
This volume gives the salient feature of the alloy. The program first utilizes the Cauchy-Gauss product, and the Cauchy-Gauss sum functions, to optimize the parameters for fitting single bands or multiple overlapping bands, also for the computation of the ordinates of the synthesized band envelope, and for an analysis of the misfit with the true spectrum. This latter program also serves as a general purpose program for subtracting spectral ordinates. The programs are written in FORTRAN IV. They are designed for card input and output in a standardized format that facilitates their interfacing with other programs developed in our laboratory for the numerical analysis of infrared and ultraviolet spectrophotometric absorption curves.
included in Volume I, Volume III, Parts 1 and 2 contain similar information for triclinic crystals.

#6500264  *4.163-3.163*
Prather, J. L.
ATOMIC ENERGY LEVELS IN CRYSTALS
NBS Monograph 19, National Bureau of Standards, Washington, D. C.
February 24, 1961
84p
Available from the Superintendent of Documents, GPO

Discrete energy levels observed within certain crystals are treated as due to perturbations of the energy levels of the free ion by an electrostatic field arising from the crystal lattice. The analytic procedures for determining the field from the charge configuration are given, and the resulting fields are classified according to their symmetry. After a general survey of group-theoretical ideas, the applicable groups are analyzed in detail, and characters appropriate for both integral and half-integral angular moments of the free ion are tabulated. These are applied to the determination of the number and type of levels arising from a free ion level with $J = \frac{5}{2}$. The results of this analysis are tabulated, as are the selection rules for electric dipole, magnetic dipole, and electric quadrupole transitions. Calculation of the perturbation matrix elements by the squares of Wigner and Racah coefficients is discussed. Examples of the application of these several techniques to specific problems are given.

#650182  *3.373*
Randall, H. M.; et al.
INFRARED DETERMINATION OF ORGANIC STRUCTURES
1949
244p
In infrared Determination of Organic Structures, methods of approach have been outlined. The possibilities inherent in the method have been explored, and currently established correlations between structure and absorption spectrum have been discussed. The instruments and techniques have been described. In Chapter III, a unique tabulation is made of the assignments of absorption bands for a large number of compounds containing double bonds. Following this, the results of several years of work on molecular structure are compiled in a form easily usable by nontheoretical chemists and physicists. Chapter V offers illustrations of the modulus graph analysis for predicting from the measured absorption spectrum to its interpretation in terms of molecular structure. Finally, the actual records of a great variety of compounds are reproduced for their reference value as a library.

#6510034  *3.273-3.273*
Boody, T. L. (Editor)
ICRP WORKING GROUP ON THERMOCHEMISTRY:
PROCEEDINGS OF THIRD MEETING MARCH 17-18, 1965,
VOLUME 1
CPIA Publication No. 92, Johns Hopkins Applied Physics Laboratory, Chemical Propulsion Information Agency, Silver Spring, Maryland
July 1965
472p
Available from DDC

This volume contains the unclassified presentations given at the Third Meeting of the Working Group on Thermochemistry, and is concerned with data of interest for rocket motor performance calculations. The 22 papers include reviews of the status of thermochemical data for the heavy element compounds and ionized species. Recent experimental progress is reported in the areas of matrix spectroscopy, mass spectroscopy, calorimetry, and high temperature vapor pressure measurements of light element compounds.

#6509001  *3.273-3.273-0.2573*
Reid, R. C.; Sherwood, T. K.
THE PROPERTIES OF GASES AND LIQUIDS:
THEIR ESTIMATION AND CORRELATION, SECOND EDITION
1966
646p
This book presents a critical review of the various estimation procedures for a limited number of properties of gases and liquids. These include critical properties, P-V-T, and thermodynamic properties, vapor pressures, latent heats, heats of formation, free energies of formation, heat capacities, surface tensions, viscosities, thermal conductivities, diffusion coefficients, and equilibrium properties of mixtures. Comparisons of experimental and estimated values are shown in the form of tables for indication of the degree of reliability of the methods discussed. The book is designed to be used as a text for molecular physics or a reference source for engineers. A set of Appendixes summarizes material referred to in the chapters and contains a large amount of tabulated data as well as references to the literature.

#6509020  *3.173*
ATOMIC MASSES FROM GALLIUM TO POLYBERÉNÉUM
Physical Review, 132: 1662-1672
November 15, 1961
The 16-inch double-focusing mass spectrometer at the University of Minnesota has been employed to measure the atomic mass of 43 stable isotopes in the region $Z = 60$ to 100. The standard error associated with these results is approximately 5 parts in $10^4$. Improvements in the instrument are described that result in an increase of useful resolution of a factor of 2 to 3. A set of radioactive masses is calculated from the stable mass data together with a beta-decay energies and nuclear reaction Q values. The resultant table of masses is used to calculate total nuclear binding energies, separation energies and pairing energies for a number of nuclei in the region near $N = 50$. The systematics of the separation energies display very smooth characteristics except at the shell closure. Neutron pairing energies show a marked decrease in value following the shell closure.

#6509025  *3.173*
Rose, R. C.
GAMMA-RAY ATTENUATION COEFFICIENTS
Nucleonics, 12:126-127
August 1954
Gamma-ray attenuation coefficients are presented for several shielding materials for gamma energies from 0.1 to 10 MeV.

#6509061  *3.273*
Barrow, R. F.; et al.
Roson, R. (Editor)
TABLES OF CONSTANTS AND NUMERICAL DATA, VOLUME S:
SELECTED TABLES OF WAVELENGTH CHARACTERISTICS OF EMISSION AND ABSORPTION BANDS OF DIATOMIC MOLECULES
Paris: Hermann & Cie, Depositaires
1952
309p
In French

This book contains tables of characteristic wavelengths of diatomic molecular bands (indicating for each band, the method of observation, the direction of degradation, the vibrational classification, etc.), the characteristic wave lengths of the bands of $C_2$, $Cl_2$, $Ar_2$, $CO_2$, $N_2$, $N_2^+$, $NH$, $NH^+$, $NO$, $O$, $OH$, $OH^+$, and the wavelengths of the radiation of $H$. A list of molecules found in parts 4 and 5 are included as well as a list of bibliographical references.
A compilation of vibrational frequency data for selected molecules is being conducted at the University of Tokyo in cooperation with the National Standard Reference Data Program of the National Bureau of Standards as a part of an international effort to compile and evaluate physical and chemical data. These tables contain fundamental vibrational frequencies of 167 molecules together with vibrational assignments, sources of data, brief comments and citations of references. The fundamental frequencies are obtained mainly from the infrared and Raman spectra. When these are not available, other experimental data such as microwave results are taken into account. The selection of vibrational fundamentals from observed spectral data is based upon careful studies of the spectral data and comprehensive mathematical analyses. These tables were designed to provide a concise summary needed for the computation of ideal gas thermodynamic properties. They may also provide a convenient source of information to those who require vibrational energy levels and related properties in molecular spectroscopy, analytical chemistry, and other fields of physics and chemistry.

**#6600018, #6990015, #6990010**
Shimanouchi, T.

**TABLES OF MOLECULAR VIBRATIONAL FREQUENCIES, PARTS 1, 2, AND 3**
March 1, 1967, October 1967, March 1968
56p, 33p, 39p

Available from the Superintendent of Documents, GPO

---

**#6602013**
Sonayajulu, G. R.; Zwolinski, B. J.

**ADDITIVITY OF THE POTENTIAL BARRIERS ABOUT DOUBLE BONDS IN PATHWAYS INVOLVING INTERNAL ROTATION**

Chemical Thermodynamic Properties Center, Texas A&M University, College Station, Texas
September 1, 1965
5p

Potential barriers about double bonds have been found to be additive of the contributions from pairs of bonds separated by C=C bond in substituted ethylenes. On the basis of this work the torsional frequency of ethylene has been found to be 325 cm⁻¹. New torsional assignments have been suggested for some molecules.

**#6610037**
Sonayajulu, G. R.; Zwolinski, B. J.

**A GENERALIZED BOND ENERGY SCHEME INCLUDING BARRIERS TO FREE ROTATION AND CORRECTIONS FOR ROTATIONAL ISOMERISM**

Chemical Thermodynamic Properties Center, Texas A&M University, College Station, Texas
1965
21p

A generalized procedure for predicting the energies of the paraffinic hydrocarbons is deduced on the assumption of (i) interactions between pairs of bonds attached to a carbon atom, (ii) interactions between torsions of bonds attached to a carbon atom and (iii) interactions between pairs of bonds separated by a C-C bond. The derived equation justifies Tatarski's empirical approach and is shown to be not very different from Allen's equation without the steric terms. Under defined conditions this equation reduces exactly to Allen's equation. Barriers to free rotation were considered in deriving the new equation. This new equation is further modified by including corrections for rotational isomerism in place of trivial interaction and the resulting equation is better founded in principle, correctly estimates enthalpies of atomization for normal and isomeric alkanes including those molecules without rotational isomers, and furthermore provides more refined steric terms due to higher order interactions.
In Part II, the enthalpies of formation of the substituted methanes have been calculated on the basis of Zahn's model. A method has been developed for the calculation of the dissociation energy of the R'-R" bond in a substituted ethane. Based on the dissociation energy of the R'-R" bond, the enthalpies of formation of the substituted methyl radicals and the substituted ethanes have been calculated. The enthalpies of formation of the substituted methyl radicals can also be calculated on the basis of Zahn's model. In Part III methods have been developed for the calculation of the magnitudes of the bond and pair bond contributions based on relationships between bond energy and bond dissociation energy. The pair bond contributions for the pair of bonds CC in CX, CX, and CX have been determined. Also, the enthalpies of formation of CX, and CS radicals, X being H, F, CI, Br, I or 'CH'. In Part IV enthalpies of formation of the CX, type radicals obtained in Part III have been used to obtain the enthalpies of formation of the CXY type radicals. Based on a relationship between bond energy and bond dissociation energy, the dissociation energies of the bond in substituted ethylenes have been calculated. Using the bond dissociation energies and the enthalpies of formation of the substituted ethylene radicals, enthalpies of formation of a few substituted ethylenes have been calculated. The calculated values have been found to be in good agreement with the observed values.

This book contains the emission spectra of 22 elements dealt with most frequently in the investigations of all forms of plasmas. For each spectral line the wavelength, intensity and classification are given. The book contains the spectra of all the gaseous elements except radon, of all the alkali metals, except rubidium and radon, the francium, radium, radon, aluminum, silicon, calcium, titanium, iron, and copper. For each element, in addition to the spectrum of the neutral atom, the spectra of several of its ions are given. The tables contain over 30,000 lines in the IR, visible, near-UV and far-UV regions of the spectrum.

The International Encyclopedia of Physical Chemistry and Chemical Physics is a comprehensive and current account of all aspects of the domain of science between chemistry and physics, written for the graduate and research worker. The subject matter is grouped in twenty topics. This third volume on the topic gas kinetics deals with energy transfer systems in gas reactions, relaxation techniques, rate measurements of competing processes, low energy level studies, and higher vibrational level energy transfer. Tables give vibrational relaxation times and collision numbers for diatomic molecules. Tables of deactivation probabilities for diatomic molecules are compiled in additive X; vibrational relaxation times, reduced relaxation times, and collision numbers for polyatomic molecules; and relative probabilities for collisional deactivation of polyatomic molecules by additives X.

The Infrared Band Handbook presents data on absorption band spectra of various organic molecules arranged by wavenumber in steps of 1 cm⁻¹ from 3610 to 617 cm⁻¹. The data for this compilation were taken from the American Petroleum Institute Project 44 series, including spectra published since 1943, and other data published between 1957 and 1963. The data included for each band are: wavenumber in cm⁻¹; intensity of the band (as: strong, medium, weak, etc.); the physical state in which the spectrum was measured or the solute used; the structural group to which the vibration was assigned in the original reference (if one was assigned) and the mode of vibration, where pertinent; and a reference number pertaining to the list of source material. There is an index to compounds arranged alphabetically by empirical formulas. The appendix contains correlation tables for methyl deformation frequencies and C-N stretching frequencies.

The interpretation of infrared spectra is based on the so-called "group frequencies" which related molecules have in common. A group frequency can be defined as a vibration which can be associated with an isolated structural group in a molecule. Each class of compounds is introduced by first presenting the vibrational analysis of a few compounds of that class, then the group frequencies found for that class, and finally the interpreted spectra. The vibrational analysis of a molecule is accomplished by calculating a series of force constants which result and lead to the observed frequencies. Substances included in Volume I are alkanes, alkenes, benzene ring
TABLES OF COULOMB WAVE FUNCTIONS, VOLUME I
NBS Applied Mathematics Series No. 17, National Bureau of Standards, Washington, D. C.
April 1952
14p
Available from Superintendent of Documents, GPO

This report presents numerical values of the Coulomb wave functions in somewhat peculiar notation. However, an introduction by M. Abramowitz defines all quantities and it is possible to reconstruct the actual Coulomb wave functions from the tables presented herein. In addition, there is a discussion of the significance of these wave functions in a preface by G. Breit.

AERODYNAMIC PHENOMENA IN STELLAR ATMOSPHERES, A BIBLIOGRAPHY
September 1959
93p
Available from Superintendent of Documents, GPO

This bibliography covers the period from 1914 to 1959. It was prepared as a working bibliography for the Fourth Symposium on Cosmic Gas Dynamics: Aerodynamic Phenomena in Stellar Atmospheres.

INFRARED SPECTROSCOPY OF CARBOHYDRATES, A REVIEW OF THE LITERATURE
NBS Monograph 110, National Bureau of Standards, Washington, D. C.
June 1968
21p
Available from Superintendent of Documents, GPO

A survey has been made of the literature on the infrared spectroscopy of carbohydrates, in order to assemble and systematize information in this field. The Monograph discusses principles and instrumentation, sampling techniques, comparison of samples, the interpretation of the spectra, particularly as regards to functional groups of carbohydrates and their derivatives, correlations for the fingerprint region and beyond, and conformational studies. A discussion, examination and discussion of the use of infrared spectra for qualitative and quantitative purposes and for the determination of structure. Special techniques are briefly described including use of plane-polarized radiation, the technique of attenuated total reflection and Raman spectra.

UV ATLAS OF ORGANIC COMPOUNDS, VOLUMES 1, 2, 3 AND 4
430p, 238p, 317p

The Atlas consists of a collection of the ultraviolet and visible spectra of compounds containing typical chromophoric groups, supplemented by tables showing the effects of substituents and solvents. The units chosen for plotting the spectra are cm⁻¹ for abscessae and a logarithmic plot of molar absorptivity. The wavenumbers covered are either 55,000 to 20,000 or 45,000 to 10,000 cm⁻¹, depending on the spectrum. The value, wave number and wavelength of maximum absorptivity are indicated explicitly on each spectrum, together with details of the compound and the conditions under which the spectrum was obtained. The sections contained in Volumes 1 to 4 include: a. compounds containing multiply bonded C atoms only; b. compounds containing C=O and C≡N; c. compounds containing multiply bonded N atoms; d. monocyclic aromatic and polycyclic aromatic; e. aromatic compounds containing non-benzenoid rings; g. monocyclic heterocyclics; h. polycyclic heterocyclics; i. derivatives of saturated organic compounds; j. selected compounds of biochemical and clinical interest; k. inorganic systems; and m. solvents. A Molecular Formula Index covers all compounds contained in Volumes 1-4.

EFFECTIVE CROSS SECTIONS FOR COLLISIONS OF ELECTRONS WITH ATOMS, ATOMIC COLLISIONS III
JILA Report 3, Joint Institute for Laboratory Astrophysics, Boulder, Colorado May 20, 1966
135p

This report consists of a series of articles on effective cross sections for collisions of electrons with atoms, as calculated theoretically by the original authors. The calculations include collisions of slow electrons with atoms of alkali metals, elastic scattering of electrons on atoms of alkali metals, excitation of neon atoms by electron bombardment, computation of excitation cross sections of excited hydrogen atoms, and computing cross sections for phases of electron-hydrogen scattering.

IONIZATION OF ATOMS BY ELECTRON BOMBARDMENT
Electron-Atom Collisions Riga 1965
100p

This is a review of the ionization of atoms by electron impact and of the theory of ionization. The article contains a bibliography of 103 references as well as 54 graphs and 5 tables of data. The article reviews the classical and quantum theory of ionization, the asymptotic behavior of the wave function, Born's Approximation, exchange calculations, calculation of distortion, momentum approximation, ionization from excited states, ionization of the helium atom and of complex atoms and ions, threshold behavior, and empirical formulas.

MICROWAVE SPECTRAL TABLES, LINE STRENGTHS OF ASYMMETRIC ROTORS
NBS Monograph 70, Volume 2, National Bureau of Standards, Washington, D. C.
December 1964
338p

Available from Superintendent of Documents, GPO

Microwave Spectral Tables, Volume 2 is the result of a computer calculation of line strengths of asymmetric rotors; the line strength is defined as a sum of squares of absolute values of direction cosine matrix elements over both laboratory-fixed Cartesian coordinates and Zeeman components. These line strengths are useful for the calculations of rotational lines in microwave and other spectra, determination of dipole moments from Stark effects or vice versa, and determination of nuclear quadrupole coupling coefficients from hyperfine splittings or vice versa. The line strengths have been tabulated as a function of the rotational quantum number J, for values of J from 0 to 35, and Ray's asymmetry parameter k. Procedures for interpolation as a function of k and extra rotation to larger J values are discussed.
For about 1500 spectral lines of diatomic molecules observed by coherent radiation techniques, measured frequencies, assigned molecular species, assigned quantum numbers, and newly computed intensities are given. Molecular data, such as rotational constants, dipole moments, and various coupling constants, determined by such techniques, are also tabulated, as are other molecular constants used in the intensity computations. References are given for all included data. For determination of hyperfine spectra, both Casimir's function and the intensity splitting factors are given for both integral and half-integral quantum numbers J and I, permitting application to additional molecules with unenclosed electronic shells and with hyperfine splitting produced by more than one nucleus.

Atomic transition probabilities for about 4,000 spectral lines of the first ten elements, based on all available literature sources, are critically compiled. The data are presented in separate tables for each element and stage of ionization. For each ion the transitions are arranged according to multiplets, supermultiplets, transition arrays, and increasing quantum numbers. Allowed and forbidden transitions are listed separately. For each line the transition probability for spontaneous emission, the absorption oscillator strength, and the line strength are given along with the spectroscopic designation, the wavelength, the statistical weights, and the energy levels of the upper and lower states. In addition, the estimated accuracy and the source are indicated. In short introductions, which precede the tables for each ion, the main justifications for the choice of the adopted data and for the accuracy rating are discussed. A general introduction contains a critical review of the major data sources.

Atomic transition probabilities for about 5,000 spectral lines of the second ten elements, based on all available literature sources, are critically compiled. The data are presented in separate tables for each element and stage of ionization. For each ion the transitions are arranged according to multiplets, supermultiplets, transition arrays, and increasing quantum emission; the absorption oscillator strength, and the line strength are given along with the spectroscopic designation, the wavelength, the statistical weights, and the energy levels of the upper and lower states. In addition, the estimated accuracy and the source are indicated. In short introductions, which precede the tables for each ion, the main justifications for the choice of the adopted data and for the accuracy rating are discussed. A general introduction contains a critical review of the major data sources.
These tables contain data helpful in the use of an x-ray spectrometer. The first section includes tables of the Bragg angles of various crystals for the first five orders of reflection, arranged by increasing angles. The data included in this section are: excited element, x-ray wavelength, order of reflection and Bragg angle for ADP, EDT, quartz 6.68, silicon, sodium chloride, gypsum, PET, LiF, quartz 2.75, and topaz. The second section includes a table of Bragg angles for each crystal mentioned above, arranged by elements in order of increasing atomic number.

About 550 references are contained in this bibliography which includes, it is believed, a complete coverage of the soft x-ray literature since 1950 and through 1960. Some references to earlier work are given, but the general review references, listed separately, give an adequate coverage of the earlier work. The emphasis is on the application of soft x-ray spectroscopy in the study of valence band electronic states in metals and alloys, and therefore, the spectral region of 25 to 800 angstroms involving ruled glass grating spectrometers is of principal interest. However, a wealth of data have been gathered, primarily by the Russians, on valence electronic states by means of high energy transitions where crystal spectrometers are satisfactory. These references and any x-ray work leading to the distribution of valence electronic states are included regardless of the transitions employed. In addition to soft x-ray data, references on all pertinent aspects of the apparatus and of experimental problems are included. Also listed separately are references of value in corroborating soft x-ray data with other results, such as energy band calculations. Subject, author, x-ray band, material, and other indices are included.

This book is a compilation of spectral lines wavelengths taken from Harrison's 1939 compilation of 100,000 lines. The first part contains the wavelengths of 43,000 lines from 60 elements, which are the most useful for identifying unknown elements. These lines are arranged by wavelength and cover wavelengths from 8,000 to 2,000 Å. The second part contains the wavelengths of 23,000 lines of 95 elements, arranged by element. The last part contains auxiliary tables, including ionization potentials, molecular weights, and melting and boiling points.
This volume is a compendium of spectral lines arranged in three parts. The first is a finding list of 349 pages in which the lines of the various spectra are arranged in the order of decreasing wavelength, from 8000 to 20000 Å. The spectrum is divided into three cases, and after each spark intensities are given. The second part contains lists of lines by element; it includes fainter lines and covers a longer spectral range than Part I. Reference source, intensity, and higher excitation potential are listed for each line. The third part contains selected tables of leading lines arranged by element, standard wavelengths, transition probabilities, and ionization potentials for the first five spectra. It concludes with a brief table giving melting and boiling temperatures for elements and some molecules.

The bibliography is presented in four principal parts: Part I is a general bibliography of 4500 entries arranged alphabetically by authors, but not including the writings of P. W. Bridgman. Part II is an index of the material of Part I, arranged by subject. Part III is a complete bibliography of the writings of P. W. Bridgman. Part IV is an index of the material in Part III, arranged by subject.

This volume contains infrared spectra of organic compounds collected from various contributing laboratories by the staff of the Thermodynamic Research Center at Texas A&M University. It contains 451 spectra presented as a plot of absorbance versus wavelength, together with the compound name, structural formula, state, temperature of measurement (concentration, and the instrument used in taking the spectrum.

This volume contains mass spectral data for 223 organic compounds presented in tabular form, with the relative intensity for each value of the mass/charge ratio tabulated for by authors, but not of the ionizing voltage. Peaks are labeled and the type of peak given. Other information given include molecular weight, formula, boiling point, structure, source, parts spectra, type mass spectrometer used, ionizing current, purity, relative and total intensity, sensitivities and temperatures of operation.

These two volumes contain nuclear magnetic resonance spectra of organic compounds collected by the staff of the Thermodynamics Research Center at Texas A&M University. They contain 622 spectra, none of which are hydrocarbons. The spectra are presented graphically, each on one sheet. Information on the compound (name, chemical and structural formula, state, solvent and temperature) and the instrument type and the frequency used are also included. The set of volumes contains a numerical index and an index of compounds.
spectroscopic data. Included are a compound name index and a formula index.

#6500200-6500205 *3.393
Zwolinski, B. J.
SELECTED INFRARED SPECTRAL DATA, SERIAL NOS. 13-2791
Chemical Thermodynamic Properties Research Center, Texas A&M University, College Station, Texas
October 1967
3000 loose-leaf pages

"Infrared Spectral Data," Volumes 1 to 8 contain infrared spectra contributed by 33 laboratories, including government, industrial, and university laboratories, which measured these spectra. Over 2600 spectra are included, nearly all are of organic compounds. Each spectrum appears on a separate page, together with the compound name, structural formula, state, temperature of measurement, and concentration. The instrument used and the laboratory taking the spectrum are also given.

#6500193-6500199 *3.393
Zwolinski, B. J.
SELECTED MASS SPECTRAL DATA
Chemical Thermodynamic Properties Research Center, Texas A&M University, College Station, Texas
April 1966
2300 loose-leaf pages

This collection of mass spectral data was compiled by the American Petroleum Institute Research Project 44 staff from mass spectra obtained experimentally from 16 laboratories. A description of the instrument and procedures used by each laboratory is given. Each mass spectrum is presented in tabular form, with the relative intensity for each value of the mass/charge ratio tabulated for different values of the ionizing voltage. Peaks are labelled and the type of peak given. Additional information concerning metastable ion transition and relative intensities for a n-butane are given, as well as data on the spectrometer sensitivity, current, voltages and temperatures of operation. Six volumes contain 1933 spectra. Volume 7 contains 154 spectra presented in matrix to the catalog.

#6500209, 6500299 *3.393
Zwolinski, B. J.
SELECTED NUCLEAR MAGNETIC RESONANCE DATA SERIAL NOS. 1 TO 594, 1c TO 124c
Chemical Thermodynamic Properties Research Center, Texas A&M University, College Station, Texas
April 1968
900 loose-leaf pages

These volumes contain nuclear magnetic resonance spectra of organic compounds collected by the staff of the Thermodynamics Research Center at Texas A&M University. The two volumes prepared for the American Petroleum Institute Project 44 contain about 600 spectra, largely of hydrocarbons. The spectra are presented graphically, each on one sheet. Information on the compound (name, chemical and structural formula, state, solvent and temperature), the instrument type and the frequency used are also included. Each set of volumes contains a numerical index and an index of compounds.

#6500210-6500211 *3.393
Zwolinski, B. J.
SELECTED RAMAN SPECTRAL DATA, SERIAL NOS. 1-400
Chemical Thermodynamic Properties Research Center, Texas A&M University, College Station, Texas
October 1964
520 loose-leaf pages

These volumes contain Raman spectra collected by the staff of the Thermodynamics Properties Center at Texas A&M University from the contributing laboratories that measured those spectra for the American Petroleum Institute Research Project 44. The line spectra are presented in tabular and graphical form with intensity of the line plotted or tabulated against the Raman shift in cm⁻¹. The intensity scale is arbitrary, with the intensity of the strongest line set equal to 100.

#6500212-6500213 *3.393
Zwolinski, B. J.
SELECTED ULTRAVIOLET SPECTRAL DATA, SERIAL NOS. 1-1075
Chemical Thermodynamic Properties Research Center, Texas A&M University, College Station, Texas
April 1967
1300 loose-leaf pages

These volumes contain ultraviolet spectra of organic compounds collected from various contributing laboratories by the staff of the Thermodynamic Research Center at Texas A&M University. The two volumes prepared for the American Petroleum Institute, Project 44 contain about 1,000 spectra. Each spectrum is presented as a plot of absorbance vs. wavelength over a range of wavelengths from 2000 to 4500Å. Also presented with each spectrum are the structural and chemical formulas for each compound, the state of the compound and the measurement temperature.
Solid State Properties
This monograph deals with the semiconducting compounds formed from elements of groups II, IV, or V and sulfur, selenium, or tellurium. The monograph describes the crystal structure of these compounds, and presents the published data on their polymorphic transitions, including transitions observed at high pressures. The monograph includes published P-T-X diagrams for the compounds discussed and describes a method for plotting the diagrams. The more important physicochemical properties of the compounds are also given: the forbidden band width, the carrier mobility, the effective masses of carriers, the electrical conductivity, and the thermoelectric power. The monograph consists of three chapters. The first deals with chalcogenides of elements in group II: zinc, cadmium, and mercury. The second discusses semiconducting compounds of elements in group IV: germanium, tin, and lead. And the third describes investigations of compounds of elements of group V: arsenic, antimony, and bismuth. The monograph does not cover chalcogenides of elements in group III. Also excluded are data on semiconductor glasses.

These books are the first two volumes of a reference source containing information on phase diagrams, crystal structure and physical and chemical properties of binary metallic systems. The first volume includes phase diagrams of systems constituted by actinium, aluminum, americium, barium, beryllium, boron, and nitrogen; the second volume includes phase diagrams of the systems constituted by bismuth, dysprosium, europium, gadolinium, gallium, germanium, hafnium, holmium, hydrogen, iron, tungsten, and vanadium. The second constituent is each element of the periodic system (except for noble gases) in alphabetical order. Subject to the extent of the present knowledge of each system, the following data are given: 1) phase diagrams and crystal structure; 2) physical properties (heat of melting or heat of mixing, surface tension and viscosity, boiling point, density, mechanical properties, heat expansion, heat capacity, thermal conductivity, electrical resistance and its temperature coefficient, magnetic properties); 3) chemical properties (resistance to corrosion under atmospheric conditions, and in various corrosive media and in gases); 4) a list of literature references.

This is a computer printout compilation of critically evaluated data, giving properties of polycrystalline aggregates of minerals containing 74 elements and anions. Literature sources of each elastic constant used in calculating properties are given. The properties tabulated for each entry include: crystalline structure; composition; density; elastic, bulk, shear, Young, and Long moduli; the Poisson ratio; and the Debye temperature.

In this supplement, additional data are compiled on the optical materials, which are useful for infrared instrumentation, treated in 2389-11-S, as well as data on three new materials, Irtran-1, Irtran-2, and diamond. In addition, several errors and ambiguities in the original report are corrected and clarified.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition, and differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing stream, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warm-up data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions
of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the optical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

Bearden, J. A.
X-RAY WAVELENGTHS
NYO-10586, Atomic Energy Commission, Oak Ridge, Tennessee
1966
53p
Available from CPSTI

This book is the result of a four-year survey by the X-ray Laboratory at the Johns Hopkins University. The primary objectives are to: (1) recompute all published wavelengths on a consistent basis; (2) include data published subsequent to the Sandstroem survey of 1957 and, in particular, reference wavelengths measured in the Johns Hopkins Laboratory; (3) convert these wavelengths to an absolute basis; (4) tabulate the data on which the recomputed values are based; and (5) include explicitly probable errors for all adopted wavelength values. The W kalpha line peak was taken as a wavelength standard. The first part of the recomputed data includes wavelength, wavelength energy values arranged according to series and by element. The K, L, M, N and O series, as well as absorption edges are given. The second part of the tables presents wavelength and energy values in order of increasing wavelength. The appendix contains an evaluation of wavelength data and presents for each line a literature reference, a published adjusted value and reason for making the adjustment, if any adjustment was made.

Bearden, J. A.
X-RAY WAVELENGTHS AND X-RAY ATOMIC ENERGY LEVELS
NSRDS-NBS-14, National Bureau of Standards, Washington, D. C.
September 25, 1967
66p
Available from Superintendent of Documents, GPO

This publication contains two critical compilations: 1) X-Ray Wavelengths. Inconsistencies in accepted values (in x units) of x-ray reference lines have recently been demonstrated, although all are supposedly based on "good" calcite crystals. Factors supporting the selection of the W kalpha line as the x-ray wavelength standard are critically discussed. A review is given of the experimental measurements which are used to establish the wavelength of this line on an absolute angstrom basis. Its value is W kalpha = 0.2090100 + 5 ppm. This may be used to define a new unit, denoted by A*, such that the W kalpha wavelength is exactly 0.2090100A*; hence lA*/lA + 5 ppm. The wavelengths of the Ag kalpha, Cu kalpha, and K kalpha have also been established as secondary standards with probable error of approximately one part per million. Sixty-one additional x-ray lines have been used as references in a comprehensive reevaluation of more than 2700 emission and absorption wavelengths. The recommended wavelength values are listed in A* units together with probable errors; corresponding energies are also given in keV. A second table lists the wavelengths in numerical order, and likewise includes their energies in keV. 2) Reevaluation of X-Ray Atomic Energy Levels. All of the x-ray emission wavelengths have recently been reevaluated and placed on a consistent A* scale. For most elements these data give a highly overdetermined set of equations for energy level differences, which were solved by least-squares adjustment for each case. This procedure makes "best" use of all x-ray wavelength data, and also permits calculation of the probable error for each energy difference. Photoelectron measurements of absolute energy levels are more precise than x-ray absorption edge data. These have been used to establish the absolute scale for eighty-one elements and, in many cases, to provide additional energy level difference data. The x-ray absorption wavelengths were used for eight elements and ionization measurements for two; the remaining five were interpolated by a Moseley diagram involving the outputs of energy levels from adjacent elements. Probable errors are listed on an absolute energy basis. In the original source of the present data, a table of energy levels in Rydberg units is given. Difference tables in volts, Rydberg, and milli-A* wavelength units, with the respective probable errors, are also included there.

Bearden, J. A.; Burr, A. F.
ATOMIC ENERGY LEVELS
NYO-2543-1, Final Report, Atomic Energy Commission, Oak Ridge, Tennessee
1965
244p
Available from CPSTI

This book contains the energy levels of atoms, and the differences between those levels, as computed from a set of x-ray wavelengths. The first set of tables contain values and probable errors of the energies of all occupied levels of each element, presented both in Rydbergs and in electron volts. The second set of tables contains differences between the energy levels presented in the first set, in units of Rydbergs, electron volts, and milli-angstroms.

Berezhnoi, A. S.
SILICON AND ITS BINARY SYSTEMS
New York: Consultants Bureau
1966
275p
Translated from Russian

This text, translated from Russian, is a review of the properties of silicon and its binary compounds. The text discusses the preparation of binary silicon compounds, properties and uses of H-Si and halogen-Silicon systems, systems formed by silicon with elements of Groups IA, IB, IIIA, IIIB, IVB, VB, VIB, VIII, transition metals of Groups IV, V, VI, VII, and VIII, and with metals of the iron subgroup, palladium subgroup, platinum subgroup, lanthanides and actinides. Ternary and quaternary systems are also included. There is a list of 716 references and tables of data accompany the systems discussed.

Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee
September 1968
42p

This bibliography contains 452 references of low temperature specific heat data and information for elements, compounds or alloys of all elements having atomic numbers 1 to 84, 90, 92, and 94, with the exception of the inert gases and technetium. The bibliography is accompanied by an index of elements and compounds.
This report gives data and information about the development and progress made in crystal growth in the United States for a variety of compounds. The demand, availability, location, and purity data are included in the information given. This report covers the year 1968.

Bilbington, D. S.; Danielson, G. C.
STATUS OF CRYSTAL GROWTH IN THE UNITED STATES
ORNL-RMIC-10, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1968
11p
Available from CFSTI

EXPERIMENTAL EVIDENCE FOR AN ENERGY GAP IN SUPERCONDUCTORS
This paper is a critical review of evidence from (a) thermal conductivity, specific heat and critical fields, (b) nuclear resonance and relaxation, and (c) microwave and infrared studies of bulk metal and of thin films. About 80 references are included.

Blondi, M. A.; et al.
EXPERIMENTAL EVIDENCE FOR AN ENERGY GAP IN SUPERCONDUCTORS
Reviews of Modern Physics, 30: 1109-1136
October 1958

PHYSICAL PROPERTIES OF MOLECULAR CRYSTALS, LIQUIDS, AND GLASSES
This book develops a methodology enabling chemists and chemical engineers to calculate certain physical properties of condensed phases to molecular structure. The states of matter treated are molecular crystals, liquids, and glasses. The physical properties covered are P-V-T and related thermal properties, and certain transport properties. Energy of vaporization, van der Waals dimensions, molar refactivity, dipole moment data and dielectric loss are used but not discussed thoroughly. Tables 14.1 to 14.16 contain the requisite molecular structure increments. Background material presented is a guide to the literature.

Bondi, A. A.
PHYSICAL PROPERTIES OF MOLECULAR CRYSTALS, LIQUIDS, AND GLASSES
New York: John Wiley & Sons, Inc.
1968
502p

The Alloy Data Center, part of the National Standard Reference Data System, has two primary functions. One is to stimulate cooperation and coordination among the existing data centers in the area of the physical properties of well characterized alloys. The other purpose is the collection (form publications as well as private communications), evaluation, and publication of data in areas of special competence in the NBS Alloy Physics Section. Of interest to the center are metals, semimetals, intermetallic compounds, and alloys. Excluded are materials which have ill-defined constitutions and heat treatments. An automated system has been developed to meet the bibliographic needs of the center. This system is described, as well as the specific properties of interest. The system presently contains a complete annotated file on papers dealing with NMR Knight shift measurements. The soft X-ray spectroscopy compilation is being kept up to date with the same system.

Carter, R. S.; Rush, J. J. (Editors)
MOLECULAR DYNAMICS AND STRUCTURE OF SOLIDS
NAS Special Publication 201, National Bureau of Standards, Washington, D. C.
June 1969
$71p
Available from Superintendent of Documents, GPO

This volume is based on material presented at the Second Materials Research Symposium of the National Bureau of Standards, held October 16-19, 1967. It provides a review of the application of spectroscopic and diffraction techniques to the study of the structure and dynamics of molecular solids. Invited papers present and practice of the major experimental methods, including neutron and X-ray diffraction, neutron inelastic scattering, infrared and Raman spectroscopy and nuclear magnetic resonance, serve as a background for the more detailed presentation and discussion of results which follows. The topics covered in a series of invited and contributed papers include the lattice dynamics of molecular crystals, the spectroscopy and crystal structure of organic and inorganic solids and the dynamics of polymers.

Cezairiyan, A.; Touloukian, Y. S.
CORRELATION AND PREDICTION OF THERMAL CONDUCTIVITY OF METALS THROUGH THE APPLICATION OF THE PRINCIPLE OF CORRESPONDING STATES
In Advances in Thermophysical Properties at Extreme Temperatures and Pressures, S. Gratch, Editor, pp.941-942.
New York: American Society of Mechanical Engineers 1965

It is shown that the Principle of Corresponding States is applicable to thermal conductivity of metals. Two correlative relations are presented. One at low temperatures (0 < T < 1.5Tm), and another at moderate and high temperatures (0.96 < T < 3Tm), where Tm is the temperature corresponding to the thermal conductivity maxima and T is the Debye temperature. A universal relation for the reduced thermal conductivity is obtained in the low temperature region based on the solid state physical theory of metals and the proposed Principle of Corresponding States.

Clark, P. V.; Dean, C. O.
FUSED SALT MIXTURES: EUTECTIC COMPOSITION AND MELTING POINTS BIBLIOGRAPHY 1907-1968
SC-R-68-106, Sandia Laboratories, Sandia Corporation, Albuquerque, New Mexico
December 1968
792p
Available from CFSTI

Data on eutectic compositions and melting points of fused salt mixtures are presented in two complementary tables arranged by systems and by melting points. Over 5,000 references are given through December 30, 1968.

Clark, P. V.; Dean, C. O.
FUSED SALT MIXTURES: EUTECTIC COMPOSITION AND MELTING POINTS BIBLIOGRAPHY 1907-1968
SC-R-68-106, Sandia Laboratories, Sandia Corporation, Albuquerque, New Mexico
December 1968
792p
Available from CFSTI

The Alloy Data Center, part of the National Standard Reference Data System, has two primary functions. One is to stimulate cooperation and coordination among the existing data centers in the area of the physical properties of well characterized alloys. The other purpose is the collection (form publications as well as private communications), evaluation, and publication of data in areas of special competence in the NBS Alloy Physics Section. Of interest to the center are metals, semimetals, intermetallic compounds, and alloys. Excluded are materials which have ill-defined constitutions and heat treatments. An automated system has been developed to meet the bibliographic needs of the center. This system is described, as well as the specific properties of interest. The system presently contains a complete annotated file on papers dealing with NMR Knight shift measurements. The soft X-ray spectroscopy compilation is being kept up to date with the same system.

Carter, R. S.; Rush, J. J. (Editors)
MOLECULAR DYNAMICS AND STRUCTURE OF SOLIDS
NAS Special Publication 201, National Bureau of Standards, Washington, D. C.
June 1969
$71p
Available from Superintendent of Documents, GPO

This volume is based on material presented at the Second Materials Research Symposium of the National Bureau of Standards, held October 16-19, 1967. It provides a review of the application of spectroscopic and diffraction techniques to the study of the structure and dynamics of molecular solids. Invited papers present and practice of the major experimental methods, including neutron and X-ray diffraction, neutron inelastic scattering, infrared and Raman spectroscopy and nuclear magnetic resonance, serve as a background for the more detailed presentation and discussion of results which follows. The topics covered in a series of invited and contributed papers include the lattice dynamics of molecular crystals, the spectroscopy and crystal structure of organic and inorganic solids and the dynamics of polymers.

Cezairiyan, A.; Touloukian, Y. S.
CORRELATION AND PREDICTION OF THERMAL CONDUCTIVITY OF METALS THROUGH THE APPLICATION OF THE PRINCIPLE OF CORRESPONDING STATES
In Advances in Thermophysical Properties at Extreme Temperatures and Pressures, S. Gratch, Editor, pp.941-942.
New York: American Society of Mechanical Engineers 1965

It is shown that the Principle of Corresponding States is applicable to thermal conductivity of metals. Two correlative relations are presented. One at low temperatures (0 < T < 1.5Tm), and another at moderate and high temperatures (0.96 < T < 3Tm), where Tm is the temperature corresponding to the thermal conductivity maxima and T is the Debye temperature. A universal relation for the reduced thermal conductivity is obtained in the low temperature region based on the solid state physical theory of metals and the proposed Principle of Corresponding States.

Clark, P. V.; Dean, C. O.
FUSED SALT MIXTURES: EUTECTIC COMPOSITION AND MELTING POINTS BIBLIOGRAPHY 1907-1968
SC-R-68-106, Sandia Laboratories, Sandia Corporation, Albuquerque, New Mexico
December 1968
792p
Available from CFSTI

Data on eutectic compositions and melting points of fused salt mixtures are presented in two complementary tables arranged by systems and by melting points. Over 5,000 references are given through December 30, 1968.
COPPER WIRE TABLES
NBS Handbook 100, National Bureau of Standards,
Washington, D. C.
February 21, 1966
41p
#6605019
Available from Superintendent of Documents, GPO

This Handbook is a revision of the Copper Wire Tables previously published as NBS Circular 31. It reflects changes in the nominal diameters of gages 45 and smaller and extends the tables to 56 gage. The Handbook provides data for the weight and length per ohm, weight per unit length, length per unit weight of copper wire, resistivity, temperature coefficient, density, and conductivity of copper wire from 0000 to 56 gage as a function of temperature from 0 to 200 °C.

THERMAL EXPANSION OF TECHNICAL SOLIDS AT LOW TEMPERATURES, A COMPILATION FROM THE LITERATURE
May 19, 1961
22p, 2p
#6600304
Available from Superintendent of Documents, GPO

Tables are given of the linear contraction relative to 293°K (L293 - L1) and the linear expansion coefficient dL/L293, of thirty elements, forty-five alloys, twenty-two other inorganic substances and twenty plastics and elastomers in the temperature range, 0 to 300°K.

THERMAL EXPANSION AND THERMAL RESISTIVITY OF METALS AND ALLOYS FROM CRYOGENIC TO ELEVATED TEMPERATURES
April 1961
206p
#6500476
Available from Superintendent of Documents, GPO

This report is a compilation of the physical properties of the metals aluminum, cobalt, iron, magnesium, nickel, platinum, and some of their alloys. The properties included are thermal conductivity, linear thermal expansion, specific heat, electrical resistivity, density, emissivity, diffusivity and magnetic permeability. Data are compiled as a function of temperature, where possible, over the range -457°F to 4500°F. The data contained in this report come principally from data sheets supplied by companies and organizations who responded to an ASTM-ASME survey, trade bulletins, published technical literature and U.S. Government publications.

BIBLIOGRAPHICAL SURVEY OF SOLID ADSORBENTS 1943 TO 1953
NBS Circular 566, National Bureau of Standards, Washington, D. C.
March 1956
152p
#6600603
Available from Superintendent of Documents, GPO

This book contains abstracts to 13763 references which give data and information on solid adsorbents published primarily between 1943 and 1953. Some references to publications before 1943 have been added from the fields of heterogeneous catalysis and surface reactions on textile materials. The coverage is restricted to heterogeneous phenomena at solid-liquid and solid-gas interfaces. The patent literature is not covered. The book is divided into chapters on the general topics of adsorption of gases and vapors on solid adsorbents, adsorption from solutions on solid adsorbents, thermal effects in adsorption processes, theories of adsorption, refining of sugar and other applications of adsorbents, general information on adsorbents and special methods of investigation, and preparation of adsorbents. A subject index, author index, and list of commercial solid adsorbents are included.

CRYSTAL DATA, DETERMINATIVE TABLES, SECOND EDITION
ACA Monograph 5, American Crystallographic Association, Oak Ridge, Tennessee
April 1, 1963
130p
#6500269
Available from Polycrystal Book Service, Brooklyn, New York

This volume presents a compilation of crystal structure data taken from literature published 1912-1960. The data can be used for identifying an unknown substance whose lattice constants have been measured or for finding structural data of a known compound. The tables are arranged by crystal class, and by axial ratio within each class (by cell length for cubic crystals). The following data are presented for each compound: axial ratios, name and chemical formula, lattice dimensions and angles (where appropriate), number of formula units per unit cell, space group, structure information, and measured and calculated specific gravity. For some compounds, melting point, color, cleavage, and optical properties are also included. The reference to the literature for each compound includes the author's name, journal name, volume, pages, and data. Both inorganic and organic compounds have been included. Indexes arranged by chemical formula and name have been included.

REPORT ON PHYSICAL PROPERTIES OF METALS AND ALLOYS FROM CRYOGENIC TO ELEVATED TEMPERATURES
April 1961
206p
#6500476
Available from Superintendent of Documents, GPO

This report is a compilation of the physical properties of the metals aluminum, cobalt, iron, magnesium, nickel, platinum, and some of their alloys. The properties included are thermal conductivity, linear thermal expansion, specific heat, electrical resistivity, density, emissivity, diffusivity and magnetic permeability. Data are compiled as a function of temperature, where possible, over the range -457°F to 4500°F. The data contained in this report come principally from data sheets supplied by companies and organizations who responded to an ASTM-ASME survey, trade bulletins, published technical literature and U.S. Government publications.

BIBLIOGRAPHICAL SURVEY OF SOLID ADSORBENTS 1943 TO 1953
NBS Circular 566, National Bureau of Standards, Washington, D. C.
March 1956
152p
#6600603
Available from Superintendent of Documents, GPO

This book contains abstracts to 13763 references which give data and information on solid adsorbents published primarily between 1943 and 1953. Some references to publications before 1943 have been added from the fields of heterogeneous catalysis and surface reactions on textile materials. The coverage is restricted to heterogeneous phenomena at solid-liquid and solid-gas interfaces. The patent literature is not covered. The book is divided into chapters on the general topics of adsorption of gases and vapors on solid adsorbents, adsorption from solutions on solid adsorbents, thermal effects in adsorption processes, theories of adsorption, refining of sugar and other applications of adsorbents, general information on adsorbents and special methods of investigation, and preparation of adsorbents. A subject index, author index, and list of commercial solid adsorbents are included.

CRYSTAL DATA, DETERMINATIVE TABLES, SECOND EDITION
ACA Monograph 5, American Crystallographic Association, Oak Ridge, Tennessee
April 1, 1963
130p
#6500269
Available from Polycrystal Book Service, Brooklyn, New York

This volume presents a compilation of crystal structure data taken from literature published 1912-1960. The data can be used for identifying an unknown substance whose lattice constants have been measured or for finding structural data of a known compound. The tables are arranged by crystal class, and by axial ratio within each class (by cell length for cubic crystals). The following data are presented for each compound: axial ratios, name and chemical formula, lattice dimensions and angles (where appropriate), number of formula units per unit cell, space group, structure information, and measured and calculated specific gravity. For some compounds, melting point, color, cleavage, and optical properties are also included. The reference to the literature for each compound includes the author's name, journal name, volume, pages, and data. Both inorganic and organic compounds have been included. Indexes arranged by chemical formula and name have been included.
paramagnetism. These materials include the elements, binary alloys, mineral compounds, double oxides and hydrates, oxides and hydrides, acids of mineral oxides, organic acids, anhydrous single and double salts, hydrated single and double salts, complex radicals, and other related data. The study of paramagnetic relaxation gives information both on the interactions between paramagnetic ions and on the interactions between the ions and the crystalline lattice. These data give magnetic techniques required for the production of very low temperatures and paramagnetic substances, Cross references and bibliographical references are included.

#6607035
Fomenko, V.S.
Samsonov, G.V. (Editor)
HANDBOOK OF THERMODYNAMIC PROPERTIES, ELECTRONIC WORK FUNCTION AND RICHARDSON CONSTANTS OF ELEMENTS AND COMPOUNDS
New York: Plenum Press Data Division
1966
151p

The experimental and theoretical calculation of the work functions of particular materials (elements and compounds) are done by various methods: thermionic, photoelectric, field-emission, effusion, contact potential difference, calorimetric and empirical relationships linking the work function to various physical characteristics which are utilized in the theoretical computation of the work function. Richardson constants are also given and references are listed at the end of the book.

#6607008
Fulkerson, W.; et al.
THERMAL CONDUCTIVITY, ELECTRICAL RESISTIVITY, AND SEEBECK COEFFICIENT OF SILICON FROM 100 TO 1300°K
Physical Review, 167:765-782
March 15, 1968

Results are presented of measurements of the thermal conductivity (90-1328 K), the electrical resistivity (300-1273 K), and the Seebeck coefficient (350-1273 K) for single-crystal and large-grained polycrystalline specimens of 99.99% pure silicon. The thermal conductivity above 387 K was measured by an absolute radial-heat-flow technique; below 350 K, by an absolute longitudinal technique. Some intermediate thermal-conductivity measurements from 300-400 K were made on the polycrystalline material using a comparative longitudinal-heat-flow apparatus. The estimated errors of these three thermal-conductivity methods were ±5 ±2%, ±1.2 ±0.1 K, and 14.0 ±9 K, respectively. The estimated error for the electrical-resistivity measurements was ±1.4 ±2 K, and for the Seebeck measurements ±1.6 ±2 K. The thermal-conductivity values were compared with conflicting data from the literatere, and they corroborate the higher-temperature results obtained by Glassbrenner and Slack. Therefore, we agree with their conclusion that the electronic contribution is reasonably close to theoretical estimates which include a large ambipolar-diffusion term. The temperature dependence of the lattice thermal resistance has been compared to various theoretical models but no approach seems to explain the data in detail. An abrupt slope change in the thermal resistivity at about 670 K is a major cause of the difficulty.

#6512006
French, J.C.
BIBLIOGRAPHY ON THE MEASUREMENT OF BULK RESISTIVITY OF SEMICONDUCTOR MATERIALS FOR ELECTRON DEVICES
October 21, 1964
118p
Available from Superintendent of Documents, GPO

This bibliography contains the references important to the measurement of bulk resistivity of semiconductor materials for electron devices. The publication contains a subject index, an index to authors, and a list of references. The subject index is divided into thirteen categories. An abstract is included for each item listed.

#6513036
SOLID STATE DIFFUSION IN METALS AND ALLOYS
AEC-tr-6315, AEC Division of Technical Information Extension, Oak Ridge, Tennessee
April 1964
52p
Available from the CFSTI Translated from the Russian

This book discusses the progress made from 1954-1958 in the investigation of solid state diffusion in metals and alloys. It describes the investigation of diffusion using radioactive isotopes and reviews interatomic interaction and laws established for diffusion in solid phase. A majority of the work presented was developed in the USSR. An extensive list of references is included as well as tables of diffusion properties. The volume includes information on the theory of diffusion, laws of diffusion and calculation for formulas, methods of investigating diffusion in metals, diffusion and interatomic interaction in alloys, defects in the crystal structure and diffusion in metals and alloys, diffusion and resistance to heat, and the use of diffusion phenomena in certain fields of technology.

#650477-6500481 (inclusive)
Goldsmith, A.; Waterman, T.E.; Hirshorn, H.J.
HANDBOOK OF THERMOPHYSICAL PROPERTIES OF SOLID MATERIALS, REVISED EDITION,
VOLUME I: ELEMENTS,
VOLUME II: ALLOYS,
VOLUME III: CERAMICS,
VOLUME IV: CERMETS, INTERMETALLICS, POLYMERIC, AND COMPOSITES,
VOLUME V: APPENDIX
New York: The Macmillan Company
1961
I:75p; II:1270p; III:1162p; IV:798p; V:286p

This set of books is an evaluated compilation of all data on the thermophysical properties of solid materials published during the period 1940-1957. Materials with melting points above 1000°F are included. The following physical properties are included: density, melting point, latent heat of fusion, latent heat of vaporization, latent heat of sublimation, specific heat, thermal conductivity, thermal diffusivity, emissivity, reflectivity, linear thermal expansion, vapor pressure and electric resistivity. Volume I contains data on the elements; Volume II, alloys; Volume III, ceramics; Volume IV, cermets, intermetallics, polymeric materials, and composite materials. Volume V, the Appendix, contains the Materials Index, the Author Index, and an alphabetic list of materials. Wherever possible, a recommended value is given.
This bibliography on pure aluminum and dilute aluminum alloys contains references on the following select properties: macroscopic structure, energy band structure, electrical resistivity, emission properties, magnetic properties, optical properties, thermal properties, Debye temperature and attenuation properties.

These data sheets present a compilation of the electrical and electronic properties of vanadium-silicon compounds, especially V-51. These properties include transition temperature, specific heat and Debye temperature, critical field, electrical resistivity, penetration depth and absorption. Other properties included are: susceptibility, energy gap, magnetic, hysterisis and susceptibility, critical current and current density, Hall coefficient and thermoelectric power. Three additional sections of this report cover the crystallographic and mechanical properties of V-51 as well as the nature of magnetic flux characteristics in the mixed state. Each of these properties is compiled over the widest possible range of parameters from references obtained in a thorough literature search.

This volume provides an updated index to the literature on the electrical and electronic properties of materials. The documents listed in the bibliography and the index represent the acquisitions of the Electronic Properties Information Center since January 1965, and the search of 63 journals and the use of eight abstracting services. Part 1 is alphabetically arranged by the name of a material, then by property. For each property, one or more accession numbers referring to a publication (if data on that material and property has been published) is listed. The specific properties tabulated are: general, absorption cross section, Curie constant, Curie temperature, Debye temperature, dielectric constant, domain structure, effective mass, electrical conductivity, electrical resistivity, electronic properties, electron field emission, electromechanical properties, electron photoemission, electron secondary emission, electron specific heat, energy bands, energy gap, energy levels, flux characteristics, gyromagnetic properties, Hall coefficient, insulation resistance, irradiation properties, lifetime, magnetic hysteresis, magnetic susceptibility, magneto-electric properties, magneto-mechanical properties, microwave emission, mobility, photoelectron properties, photon electroluminescence, photon emissivity, photon transmission, reflection coefficient, piezoelectric properties, refractive index, superconductor transition temperature, thermal conductivity, thermoelectric properties, thermomechanical properties, transport properties, and work function. Part 2 contains the bibliography arranged by accession number. For each publication there is listed the author's name, title of the publication, name, volume, date and page number of the journal.

These data sheets present a compilation of a wide range of electrical and magnetic properties for nickel-tungsten alloys of various purities in bulk and film form. Electrical properties include conductivity, resistivity, Hall coefficient and transition temperature. Magnetic properties include susceptibility, hysterisis and threshold field. Properties and data include thermal, magnetic, photoelectric properties, photon emissivities, absorption and energy gap. Debye temperature and thermal conductivity data are also given. Each property is compiled over the widest possible range of parameters from references obtained in a thorough literature search.

These data sheets, as Part I of two parts, represent a compilation of the structure-sensitive properties of niobium-tin composites. These properties include thermal conductivity, critical current and current density and solenoid device applications. Finally, the properties of niobium-tin ternary compounds are presented. Each of these properties is compiled over the widest range of parameters from the references obtained in a thorough literature search.

These data sheets, as Part II of two parts, represent a compilation of the structurally insensitive properties of niobium-tin composites. These properties include phase diagrams, crystallographic properties, specific heat, and transition temperature, magnetic properties, penetration depth and electrical resistivity. Each of these properties is compiled over the widest range of parameters from the references obtained in a thorough literature search.

These data sheets present a compilation of the electrical and electronic properties on niobium-zirconium alloys. These properties include transition temperature, specific heat and Debye temperature, resistivity, bulk and film conductivity, electrical resistivity, electrical and thermal conductivities and ultrasonic attenuation. Other properties included are: superconducting energy gap, magnetic hysteresis and susceptibility, critical current and current density, bulk and film voltage and thermoelectric properties.
power. Three additional sections of this report cover the crystallographic and mechanical properties of Nb-Zr alloys as well as the nature of magnetic flux characteristics in the mixed state. The use of these alloys in solenoid applications is also discussed. Each of these properties is compiled over the widest possible range of parameters from references obtained in a thorough literature search.

#6500653 *4.133-3.133-8.133
GROUPS IV, V, AND VI TRANSITION METALS, SELECTED REFERENCES ON PREPARATION, STRUCTURE, PHYSICAL PROPERTIES, METALLOGRAPHY, AND TRANSPORT PROPERTIES ORNL-RMIC-3 (Rev.), Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee April 1965 94p

This bibliography contains lists of selected references on preparation methods, crystal structure, transport properties, and physical and mechanical properties for the Groups IV, V, and VI transition metals. An abstract is included along with each reference entry.

#6902997 *4.133-3.133-8.133
GROUPS IV, V, AND VI TRANSITION METALS, SUPPLEMENT NO. 1, SELECTED REFERENCES ON PREPARATION STRUCTURE, PHYSICAL PROPERTIES, METALLOGRAPHY, AND TRANSPORT PROPERTIES ORNL-RMIC-8, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee April 1965 27p
Available from CFSTI

This bibliography is a supplement to ORNL-RMIC-3 (Rev.) of April 1965, and contains lists of selected references on preparation methods, crystal structure, transport properties, and physical and mechanical properties for the Groups IV, V, and VI transition metals.

#6500465 *8.133-4.133

Thermal radiation property values for more than 40 metals and their alloys are given. The properties reported are emissivity, absorptivity, reflectivity, and transmissivity at various surface conditions, temperatures, and wavelengths. In addition, radiation properties of building materials, paints, glasses, papers, cloths, and other miscellaneous solid materials have been compiled, and solar absorptivity and equilibrium temperatures for various materials have been tabulated. These values have been gathered from about 200 references in English, German, French, and Russian and have been carefully analyzed to obtain, in the author's opinion, the most trustworthy data for practical application. Recommended data for technically important materials are also presented graphically. An extensive survey was conducted to compile values for Stefan-Boltzmann's, Planck's, and Wien's constants, and the most reliable values for practical application are listed in a separate table.

#6806004 *4.133
Available from Superintendent of Documents, GPO

Experimental electrical resistivity data for 16 pure metals have been compiled, tabulated, and graphically illustrated for a temperature range of 0 to 273*K. A section has been prepared for each particular metal which includes references, brief comments concerned with preparation of sample, purity, and other pertinent information, tabulated data, and graphs. Data are presented for aluminum, beryllium, cobalt, copper, gold, indium, iron, lead, magnesium, molybdenum, nickel, niobium, platinum, silver, tantalum and tin.

#6800901 *8.233-4.233
Available from the Superintendent of Documents, GPO

The work presented in this report comprises the critical evaluation, analysis, synthesis of the available thermal conductivity data, and the generation of recommended values for twelve metallic elements, mainly for the solid state, for a range of graphites and for three fluids in the gaseous state. These are cadmium, chromium, lead, magnesium, molybdenum, nickel, niobium, tantalum, tin, titanium, zinc, zirconium, Acheson graphite, ATJ graphite, pyrolytic graphite, 875S graphite, 890S graphite aceton, ammonia and methane. For each of the materials recommended values are given over a wide range of temperature.

#6500093 *8.133-4.133

This volume contains the results of evaluations for all the metallic elements and binary alloy systems for which thermodynamic data were available. Results of the evaluations in the form of discussions and analyses of the data and tables of selected values are given for 63 elements and 168 alloy systems. Some of the properties covered include freezing point, vapor pressure, transition temperature phase diagrams, entropy, heat capacity, free energy function, heat content; heat of transition, sublimation, fusion and vaporization; entropy of transition, fusion and vaporization; free energy of formation; free energy - excess integral, relative partial molar and relative partial molar excess; heat of formation - relative partial molar; entropy of formation - excess integral, relative partial molar and relative partial molar excess; heat capacity - relative partial molar.

#6511007 *8.133-4.133
Hultgren, R.; Orr, R.L.; Kelley, K.K., SUPPLEMENT TO SELECTED VALUES OF THERMODYNAMIC PROPERTIES OF METALS AND ALLOYS California University, College of Engineering and Lawrence Radiation Laboratory, Berkeley, California June 1970 500+ loose-leaf pages

This Supplement contains the results of evaluations for 56 metallic elements and 178 binary alloy systems for which thermodynamic data were
available. Reevaluation and extension of the selected values will be a continuing process as new data become available. Prior to the publication of a revised edition of the book, additional evaluations, as they are completed, will be issued in loose-leaf form at frequent intervals for inclusion with this Supplement. Description of the methods of evaluation and forms of presenting the data may be found in the parent volume of "Selected Values of "H. R. * Reise of the evaluation in the form of discussions and analyses of the data and tables of selected values are given. Some of the properties covered include freezing point, vapor pressure, transition temperature, phase diagrams, entropy, heat capacity, free energy function, heat content, heat of transition, sublimation, vaporization; entropy of transition, fusion and vaporization; free energy of sublimation, vaporization; free energy of formation; free energy - excess integral, relative partial molar and relative partial molar excess; heat of formation - relative partial molar; entropy of formation - excess integral, relative partial molar and relative partial molar excess; heat capacity - relative partial molar.

*4,143

INSULATION MATERIALS - POLYETHYLENE TEREPTHALATE, POLYTETRAFLUOROETHYLENE PLASTICS, POLYTETRAFLUOROCHLOROETHYLENE PLASTICS, DS-105, 106, 107, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California June 1962 32p, 31p, 14p

These reports contain data sheets on the properties of plastics: polytetrafluoroethylene, polytetrafluoroethylene and polyethylene terephthalate. Each report contains data on some or all of the following properties: arc resistance, carbonization strength, the dependence on constant, dielectric strength, dissipation factor, insulation resistance, power factor, surface resistivity, and volume resistivity.

*4,143-4,143

Janssen, J.; Luck, J.; Torborg, R.

REFLECTANCE OF ANODIZED TITANIUM AND BERYLLIUM Aeronautical Systems Division, Wright-Patterson Air Force Base, Ohio (No date) 32p

Selection of materials having the desired thermal radiation properties is of primary importance for space applications, since heat transfer to and from an object in space occurs only by radiation. The effect of anodizing variables on the reflectance of anodized titanium and beryllium was studied over the range of 0.4 to 22 microns in a vacuum of 10^-5 mm Hg with specimem temperatures of 100 to 825 F. In general, the anodizing process had a greater effect on the reflectance than did alloying elements in the metal. The pressure of water in some coatings, which was driven off at elevated temperatures in the vacuum, gave lower reflectance in the infrared.

*4,243

Kennard, O.; Watson, D. C.


This bibliography of organic crystal structures covers the period 1960-1966 and provides references to about 2,000 compounds whose structures were determined by X-ray or neutron diffraction. The bibliography forms part of a computer-oriented structural Library which contains reference material, numerical data including atomic coordinates and thermal parameters, and descriptive information. The bibliography includes compound name, author and journal references.

*8,143-4,143

Kolsky, H. G.

THE THERMODYNAMIC PROPERTIES OF 54 ELEMENTS CONSIDERED AS IDEAL MONATOMIC CASES LA-2110, Los Alamos Scientific Laboratory, Los Alamos, New Mexico March 15, 1957 138p Available from CFSTI

Tables of the thermodynamic properties of fifty-four elements have been calculated considering each as an ideal monatomic gas. The energy levels as determined by the National Bureau of Standards Spectroscopy Section were prepared on IBM cards for one of the LASL IBM 704 computers. The partition functions were calculated for each element and the thermodynamic properties were evaluated from them for 150 temperatures between 10^6 K and 8000K. A short table of the harmonic oscillator functions was also done in the same format as the other tables for use in evaluating the vibrational contributions to the thermodynamic properties. A brief discussion of the theory and calculational details is given.

*4,243

Krishnan, R. S.

PROGRESS IN CRYSTAL PHYSICS, VOLUME I: THERMAL, ELASTIC AND OPTICAL PROPERTIES Madras, India: S. Viswanathan 1958 198p

This volume contains data and a bibliography on the physical properties of crystals current as of December 1957, for general reference in research on crystal physics. Seven authors review thermal expansion, thermal conductivity, elastic properties, photo-elastic properties, thermo-optic behavior, Faraday Effect and dielectric behavior of crystals. Metal crystals and polycrystalline materials are excluded. Each chapter discusses the theory, experimental methods, measurements, and results. The available data on each property for single crystals have been collected and presented in a tabular form at the end of the chapter.
This section contains data on the atomic physical interpretation of crystal physics as of November 1954. The following properties have been tabulated: the characteristics of symmetry, elements, the 32 crystal classes, space groups, lattice types, structures, crystal dimensions of inorganic and organic crystals, atomic and ionic radii, lattice energies, internal vibrations of crystals, electron emission, energy bonds, X-ray spectra and bonding states, electron spectra of crystals, and high frequency spectra of crystals.

This section of the Landolt-Bornstein tables contains data on the electrical properties of matter (except electrochemical systems) published through February 1959. The properties tabulated include the conductivities of pure metals and certain alloys at 0 °C and their temperature variations, resistance ratios, effect of magnetic fields on resistance, superconducting metals and alloys and their transition temperature, magnetic field effects on super conductivity, galvanomagnetic and thermomagnetic effects, ionic conductivity in crystals, transfer numbers in solids. Also tabulated are the constants of homogeneous semiconductors; photoelectric conduction; piezoelectric equations; and the elastic, piezoelectric and dielectric constants of piezoelectric crystals; dielectric properties of inorganic and organic crystals and crystalline fluids and pure fluids; dielectric properties at high frequencies; temperature and pressure dependence of dielectric constants; and the dielectric properties of gases. Thermionic emission and work functions, thermopotentials, Peitler and Thomson heats, and photoemission data are also listed.

This section of the Landolt-Bornstein tables contains tables and curves of magnetic data for paramagnetic, ferromagnetic, ferrimagnetic and antiferromagnetic substances, taken from the literature published through September 1962. The following quantities have been tabulated: fundamental magnetic properties (dia- and paramagnetic susceptibility, spontaneous magnetization, and Curie temperatures) for metallic elements and alloys, exchange energy, crystal energy, magnetostriiction, magnetocaloric effects, Faraday effect, magnetooptical effect, Kerr effect, gyromagnetic effects and resonance in ferromagnetic metals and alloys, properties of ferropinels, magnetic garnets, perovskites and hexagonal ferrites, magnetic properties of transition element compounds, lanthanide series compounds and actinide series compounds, paramagnetic centers in crystals, paramagnetic ions, ionization constants, magnetic susceptibility, cyclotron resonance, de Haas-Schubnikow effect and Faraday effect in semi-conductors, paramagnetic relaxation and nuclear magnetic relaxation.
This section of the Landolt-Bornstein tables gives the paramagnetic and diamagnetic properties of substances. The compiler has listed one or more values for each substance, and marked with an asterisk those considered acceptable and reliable. These, and other selected values considered worthy of inclusion, are listed under "Used References" and values far outside the others under "Further References." Flow values were taken from literature up to April 1966. Nolar magnetic susceptibilities are listed for compounds, and magnetic mass susceptibilities for elements. Diamagnetic general properties and inorganic substances, and principal susceptibilities for anisotropic substances. Paramagnetic susceptibilities are listed for organic compounds only.

This section of the Landolt-Bornstein tables contains material values and mechanical behavior of non-metallic substances, and their technical applications. Data are compiled from literature published through autumn 1964. The following general properties of non-metallic solids are given for natural and synthetic building materials, organic natural materials (wood, paper, cellulose, wood pulp), fibrous materials, ceramics, glasses, synthetic materials, and natural and synthetic materials. Also presented are data on friction and rolling resistance of non-metallic and metallic substances, viscosity, flow through tubes, circuitous flow of bodies during two-dimensional, flow and rough flow, flow, flotion, speed of sound, sound radiation and absorption, the human ear, sound spectra, vibrational numbers, spatial and building acoustics, sound recording and biological effects of ultrasonics.

This section of the Landoldt-Bornstein tables includes data on metallographic principles and concepts, testing properties, and the properties of iron and its alloys. Data are given on diagrams of state, strength testing and destruction free materials. The physical properties of iron tabulated are lattice constants, density, thermal expansion, diffusion, atomic heats, specific heats, entropies, entropy and free energies, phase diagrams and transition points, thermal conductivity, electrical properties, magnetic properties, optical constants and excitation cross-sections. Composition, treatment, strength values and special chemical properties of pure iron, unalloyed steels, highly alloyed steels, special steels, and cast iron are presented.
This section of the Landolt-Bornstein tables deals with the technological aspects of electricity, light and X-rays. It is compiled from the literature published through 1956. Wherever possible, the compilers have drawn from several values available for a given quantity, the value then given is most accurate. The properties tabulated in the electrical section are electrical conductivity, resistance and temperature coefficient of resistance; conductivity of molten metals at fixed pressure, tension and cold machining on electrical resistance; resistance as functions of frequencies; resistance of thin layer semiconductors; thermoelectric emf of metal pairs and thermocouples in use; effects of pressure and cold treatment on thermocouples; conduction of electricity in gases, breakdown potentials of gases; insulating liquids, filling and impregnating media, fibrous material insulators, insulating foils, mica insulators, rubber insulators; resin, plastics, quartz, glass, ceramic materials and rocks as insulators; and magnetic materials. The quantities tabulated in the light section are: sensitivity of the human eye, illuminating materials, light sources, light filters and photographic films. The X-ray section includes data on measurements and units of X-ray technology.

**#6804008**

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME IV: TECHNOLOGY, PART III: HEAT TECHNOLOGY, HEAT TECHNOLOGY ENGINEERING MEASURING METHODS, THERMODYNAMIC PROPERTIES OF HOMOGENOUS MATERIALS

Hausen, H. (Editor)

Berlin, New York: Springer-Verlag 1967

944p

In German

This section of Landolt-Bornstein is a compilation of data dealing with the technology of thermal measurements and the thermodynamic properties of gases, vapors, liquids, and solid materials. Data are included for such areas as the basis of temperature measurement; the temperature measurement with contact thermometers; optical temperature measurement (Pyrometry); hygrometry; thermodynamic properties of homogenous materials, gases, water and steam, refrigerating agents; thermal expansion of solids and liquids; and the specific thermal properties of technically important materials.

**#6706020**

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW SERIES, GROUP II: ATOMIC AND MOLECULAR PHYSICS, VOLUME 2: MAGNETIC PROPERTIES OF COORDINATION AND ORGANO-METALLIC TRANSITION METAL COMPOUNDS

Koenig, E.

Hellwege, K.-H.; Hellwege, A.M. (Editors)

Berlin, New York: Springer-Verlag 1966

576p

In German and English

Magnetic properties are listed for coordination and organo-metallic transition metal compounds. The magnetic susceptibility section lists for each compound gram susceptibility and molar susceptibility at a particular temperature; magnetic moments and paramagnetic Curie temperatures, and the experimental method used to obtain data. The electron paramagnetic resonance section lists for each compound the spectroscopic splitting g-factor, fine structure and hyperfine structure parameters, half-width of the EPR line, and temperature and frequency of measurement.
Volume III/2 of the New Series of Landolt-Börnstein continues Volume III/1 only three years after publication of the latter. All the tables contained in III/1 are brought up to date by supplements in III/2. In addition, the elastic and electrooptic constants of higher order and the nonlinear optical susceptibilities have now been included. Readers should note that the Index of Substances at the end of III/2 refers to both volumes.

#6511001 *4.135
LOW-TEMPERATURE PHYSICS, PROCEEDINGS OF THE NBS SEMICENTENNIAL SYMPOSIUM ON LOW-TEMPERATURE PHYSICS HELD AT THE NBS ON MARCH 27, 28 AND 29, 1951
NBS Circular 519, National Bureau of Standards, Washington, D. C.
October 6, 1952
29p
Available from Superintendent of Documents, GPO

The papers published in this book were presented at the NBS Semicentennial Symposium on Low-Temperature Physics and contain sixty-one articles representing scientific findings in low-temperature materials. The book examines the superconductivity of various elements and isotopes and the low-temperature effects on atomic heat, resistance, specific heat, electrical conductivity, Eddy currents, and supercurrents. The book includes discussions of liquid properties of gases and the solidification of gases and gives optical and spectral data of the low-temperature effects.

#6500220 *4.153
Lyman, T. (Editor)
METALS HANDBOOK, VOLUME 1: PROPERTIES AND SELECTION OF METALS, EIGHTH EDITION
ASM Metals Handbook Committee, American Society for Metals, Metal Park, Ohio 1961
1300p

This book is a comprehensive monograph covering alloy steels, aluminum alloys, carbon steels, copper alloys, magnetically soft materials, permanent magnet materials, stainless steels and tool steels. Data are presented for magnetic, electrical, and special purpose materials. Nickel and nickel alloys, tin and tin alloys, zinc and zinc alloys, and precious metals. Physical property data are included.

#6611022 *4.153
Lyman, T. (Editor)
METALS HANDBOOK, VOLUME 2: HEAT TREATING, CLEANING AND FINISHING, EIGHTH EDITION
ASM Metals Handbook Committee, American Society for Metals, Metal Park, Ohio 1964
708p

This is the second in a series of five volumes that supersede and expand the 7th edition of Metals Handbook. The aim has been to provide information helpful in selecting and controlling processes for heat treating, cleaning and finishing of ferrous and nonferrous metals. The material on heat treating is organized into eight principal sections, four of which deal largely with processes for heat treating carbon and low-alloy steels, and four of which deal with the heat treating of metals other than carbon and low-alloy steels. Articles on heat treating of stainless steel, heat-resisting alloys, and various nonferrous metals are included. The subject matter on cleaning and finishing is organized into seven principal sections, five of which deal with iron and steel, and two of which deal with other metals.

#6500418 *4.113
LaGuillaume, C.; et al.
TABLES OF CONSTANTS AND NUMERICAL DATA, VOLUME 12.
SELECTED CONSTANTS RELATIVE TO SEMICONDUCTORS
76p

In French, preface in French and English

This "Table of Selected Constants" lists constants and numerical data for materials which exhibit a semi-conducting character. It gives the semi-conducting and physicochemical properties for germanium, silicon, tellurium, selenium, and diamond and for the I - VIII compounds. For germanium, silicon and the III-V compounds, extrinsic properties are also given (activation energy, diffusion coefficient and solubility of impurity elements). The numerical data have associated references to the original work from which they were taken. The physical quantities listed for each include symmetry group and crystal parameters, refractive index, dielectric constant, effective ionic charge, work function, photoemission work function, piroelectricity, phonon and Debye temperatures, magnetic susceptibility, coefficient of linear expansion, fusion temperature, sublimation temperature, specific heat, latent heat of fusion and sublimation, thermal conductivity, disorder factor, and density. In addition to data specific to band structure, the Table also includes mobilities of electrons and their variation with temperature.

#6609015 *4.152
LISTS OF AVAILABLE AND DESIRED MATERIALS AND AVAILABLE CRYSTAL-GROWTH REFERENCES (ULTRAPURE INORGANIC RESEARCH MATERIALS)
ORNL-RMIC-5, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee August 1966
45p
Available from the CFSTI

The materials in the Available Materials list are preferably single crystals selected for highest purity and largest dimensions. Other forms (wires, powder, foils, etc.) and lower purities and smaller sizes are usually available. Those entries marked with an asterisk were produced in the course of research, and samples might or might not be available for loans or exchange. All others are commercially produced and others listed in addition to other characteristics are in most cases those claimed by the producer.

#6902095 *4.172
SUPPLEMENTAL LISTS OF AVAILABLE AND DESIRED MATERIALS AND AVAILABLE CRYSTAL-GROWTH REFERENCES (ULTRAPURE INORGANIC RESEARCH MATERIALS)
ORNL-RMIC-6, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee August 1967
13p
Available from CFSTI

The materials in this list are single crystals selected for highest purity and largest dimensions. Other forms (wires, powder, foils, etc.) and lower purities and smaller sizes are usually available. Those entries marked with an asterisk were produced in the course of research, and samples might or might not be available for loans or exchange. All others are commercially produced for sale. Purities and other characteristics are in most cases those claimed by the producer. This list is a supplement to the Available Materials List in ORNL-RMIC-5.
A compilation of the electrical properties of various borosilicate glasses is presented. Arrangement of the data sheets are by Corning code or number with a Master Identification Chart relating the known chemical composition of each glass. Detailed electrical properties include corona effects, dielectric constant, dielectric strength, dissipation factor, loss factor, power factor, surface and volume resistivity. Each property is compiled over the widest possible range of temperature and frequencies from references obtained in a thorough literature search. The structure, applications and phase diagrams are briefly reviewed.

**#680023**  
Milek, J. T.  
EPITAXIAL SILICON AND GALLIUM ARSENIDE THIN FILMS ON INSULATING CERAMIC SUBSTRATES, A STATE-OF-THE-ART REPORT  
S-9, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California August 1968  
14p  
Available from DDC, AD 675 578

A state-of-the-art literature survey is presented on the success of epitaxial deposition of silicon and gallium arsenide thin films on ceramic substrates for device applications. The electrical, thermal, chemical and crystallographic properties of sapphire, spinel, beryllium oxide, magnesium oxide, diamond, quartz, silicon carbide, aluminum silicate glass, glazed ceramic and other miscellaneous ceramic substrate materials are presented. A master flow chart is developed to identify each variable at each stage of the film/substrate preparation and resultant device fabrication processes. A series of charts identifying each of these variables as reported in the literature for each substrate material is presented. A complete bibliographic review of silicon epitaxial deposition, independent of substrate, is also presented.

**#6500435**  
Milek, J. T.  
FORSTERITE DATA SHEETS  
DS-129, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California August 1963  
28p

This report contains a compilation of the insulation properties of Forsterite (2 MgO·SiO.). The information was taken from literature published through August 1963. The following quantities are tabulated: dielectric constant, dielectric strength, electrical resistivity, loss factor, power factor and Tg value.

**#6500434**  
Milek, J. T.  
PYROCERAM DATA SHEETS  
DS-130, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California August 1963  
37p

This report contains a compilation of the insulting properties of Pyrocram, a new family of glass materials developed by Corning Glass Works. Seven types of Pyrocram are discussed; of these, however, only two were commercially available as of August 1963. The properties of these two are: dielectric constant, dielectric strength, dissipation factor, loss factor and volume resistivity.
A compilation of the electrical properties of sulfur hexafluoride, a dielectric gas, is presented. Electrical properties include corona, dielectric constant, dissipation factor and dielectric strength. The latter property data section is segregated into parameter effects as follows: pressure, gap distance, temperature, electrical configurations and gas mixtures. Each property is compiled over the widest possible range of parameters obtained in a thorough search of the world's literature.

A survey of the following properties of tetrafluoroethylene plastics was made: mechanical, electrical, thermal, chemical and irradiation behavior. Each property set of data represents the greatest range of parameters: temperature, frequency, pressure, voltage and environmental conditions (vacuum, radiation, etc.) found in the literature. The effects of fillers on every property are also detailed where such data were available. Applications such as space, mechanical, electrical, and cryogenic, are treated in some detail for the various fabricated forms of tetrafluoroethylene. A brief discussion of forming and fabrication, joining and fastening methods, and sources of specifications to aid the designer in selecting tetrafluoroethylene plastics for various applications is present.

A state-of-the-art survey and a data sheet compilation of elemental boron are presented. The thorough literature search uncovered 304 references which were evaluated for information and data on the (1) reduction preparation methods, (2) zone-refining and melting techniques, (3) crystal structures, (4) electrical properties, (5) optical properties, (6) thermal and thermodynamic properties, (7) physical and chemical properties, (8) metallurgy and mechanical properties, (9) applications, and (10) availability and suppliers. Major emphasis was the characterization of boron's semiconductor and acoustic properties with respect to potential semiconductor and delay-line devices. Detailed information on the various methods of preparing pure boron and the resulting crystal structures plus properties are summarized in tables.

All commercially available aluminum alloys are identified by means of their standard designations.
This report contains a compilation from literature published through November 1963 of the semiconductor properties of cadmium telluride. The properties of cadmium telluride considered are: absorption, cross-sections, Debye temperature, dielectric constant, effective mass, electrical conductivity, electrical resistivity, energy bands, energy gap, energy levels, Hall coefficient, irradiation effects, mobility, photoelectron properties, photon luminescence, piezoelectric properties, polarization, refractive index, thermal conductivity and thermoelectric properties.

These data sheets present a compilation of a wide range of electronic properties for cadmium sulfide. Electrical properties include conductivity, dielectric constant, Hall coefficient and mobility. Emission data have been broken down into the varied electron and photon emissions which result from application of electromagnetic energy over a wide spectrum and a wide variety of photoelectric phenomena is shown. Energy data include energy bands, energy gap and energy levels as well as effective mass tables and work function. The optical properties include absorption, reflection and refractive index. Magnetic data are presented, as well as several other physical phenomena, such as Debye temperature. Thermoelectric and thermomagnetic properties are shown. Each property is compiled over the widest possible range of parameters including bulk and film form from references obtained in a thorough literature search. A summary of crystal structure and phase transitions has been included.

These data sheets present a compilation of the semiconductor properties of cadmium telluride, taken from literature published through June 1962. The quantities tabulated include absorption, Debye temperature, dielectric constant, effective mass, electrical conductivity, emission, energy bands, energy gap, Hall coefficient, irradiation effects, magnetoelectric properties, mobility, photoelectric properties, reflection, refractive index, resistivity and thermoelectric properties.

These data sheets present a compilation over a wide range of electronic properties for cadmium telluride. These properties are compiled over the widest possible range of parameters and are then aggregated in several large groups as follows: optical properties include absorption, reflection and refraction. Transport properties include electrical conductivity, Hall coefficient, mobility, effective mass, lifetime and magnetoresistance. Energy band structure includes energy gap and levels, as well as work function values. Phonon branch distribution appears separately. Both photon and electron emissions, data, are represented. Thermal properties include Debye temperature, thermal conductivity, and thermal expansion. There are other individual properties and effects included, such as dielectric constant, piezoelectric values, irradiation effects and magnetic susceptibility. A large number of graphs illustrate the various photoelectric phenomena. A section on general properties includes crystal, phase diagrams, vapor-pressure data, and preparation of single crystals and films. A table of the best values available for all physical and electronic properties is given. A separate section is given for the cadmium telluride-mercury telluride system and device applications for both the cadmium telluride crystals and the mixed crystals are appended.

These data sheets present a compilation of a wide range of electrical, optical and energy values for pure and variously-doped germanium in bulk and film form. Electrical properties include conductivity, resistivity, dielectric constant, Hall coefficient, mobility, lifetime, thermoelectric and thermomagnetic effects. Emission data has been broken down into the varied electron and photon emissions which result from application of energy in the electromagnetic spectrum. Energy data include energy bands, energy gap and energy levels for variously-doped germanium, as well as effective mass tables, work functions, carrier diffusion and cross sections. The optical properties include absorption, reflection and refractive index. Other magnetic data and irradiation effects are included, as well as several bordering physical phenomena such as piezoelectric properties, Debye temperature and thermal conductivity. Each property is compiled over the widest possible range of parameters from references obtained in a thorough literature search. A crystallographic summary has been added.

These data sheets present a compilation over a wide range of electronic properties for cadmium telluride. These properties are compiled over the widest possible range of parameters and are then aggregated in several large groups as follows: optical properties include absorption, reflection and refraction. Transport properties include electrical conductivity, Hall coefficient, mobility, effective mass, lifetime and magnetoresistance. Energy band structure includes energy gap and levels, as well as work function values. Phonon branch distribution appears separately. Both photon and electron emissions, data, are represented. Thermal properties include Debye temperature, thermal conductivity, and thermal expansion. There are other individual properties and effects included, such as dielectric constant, piezoelectric values, irradiation effects and magnetic susceptibility. A large number of graphs illustrate the various photoelectric phenomena. A section on general properties includes crystal, phase diagrams, vapor-pressure data, and preparation of single crystals and films. A table of the best values available for all physical and electronic properties is given. A separate section is given for the cadmium telluride-mercury telluride system and device applications for both the cadmium telluride crystals and the mixed crystals are appended.
These data sheets present a compilation of a wide range of electronic properties for lead oxide. Electrical properties include conductivity, resistivity and dielectric constant. For each material, a variety of electronic phenomena are shown. Energy data include energy bands, energy gap and energy levels, as well as effective mass tables, phonon energy dispersion and work functions. The optical properties include absorption, reflection and refractive index. Data on several physical phenomena, such as thermal conductivity, Debye temperature, magnetic susceptibility, and Richardson's Constant are presented. Thermoelectric data is given. Each property is compiled over the widest possible range of parameters including bulk and film form, from references obtained in a thorough literature search. A summary of crystal structure and phase transitions has been included.

This report contains data sheets on magnesium stannide compiled from the literature, giving values for absorption, effective mass, electrical conductivity, energy gap, energy level, Hall coefficient, magnetoelastic properties, magnetic susceptibility, mobility, photoemission properties, resistivity, thermal conductivity and thermoelectric properties. This compilation deals only with magnesium stannide as a semiconductor. A list of references to the literature is included. Data are given graphically and in tabular form.

These tables include the most reliable information available to date for every property -- mechanical, crystallographic, thermal, magnetic, electronic and optical -- of each of the 2-6 binary semiconducting compounds.

This interim report represents a supplement to the Electronic Properties Information Center Data Sheets on Silicon Carbide, published in June 1955 (DS-145 by M. Neuberger, AD 465 161). Almost 100 new documents concerned with silicon carbide are listed. Selected graphs and/or tables are presented from those articles containing data of particular interest. A data table of physical and electronic property values has been added to allow for a rapid review of the important properties of the alpha- and heta- crystalline forms of silicon carbide.

This report is a compilation of the semi-conductor properties of zinc oxide. The information was compiled from literature through October 1963. The report contains data on: absorption, Debye temperature, dielectric constant, effective mass, electrical conductivity, electrical resistivity, energy bands, energy gap, energy levels, Hall coefficient, irradiation effects, lifetime, magnetoelectric properties, photoelectric properties, photoelectron properties, photon luminescence, phonon thermoluminescence, piezoelectric properties, reflection coefficient, thermal conductivity, thermoelectric properties, and work function.

This report is a compilation of the semi-conductor properties of zinc selenide. The information was compiled from literature published through September 1963. The report contains data on: absorption, Debye temperature, dielectric constant, effective mass, electrical conductivity, electrical resistivity, electroacoustic properties, energy bands, energy gap, energy levels, mobility, photoelectric properties, photon luminescence, piezoelectric properties, reflectivity, refractivity, index and thermal conductivity.

A wide range of electrical and optical properties are covered in these data sheets on magnesium oxide. Particular emphasis has been placed on the dielectric properties and the energy band structure. A data table containing the best available information on the crystallographic, thermal, mechanical, physical, electronic and photoemissive properties is included.

This report contains a compilation of the semi-conductor properties of zinc sulfide. The information was compiled from literature published through December 1963. The report contains data on: absorption, cross sections, Debye temperature, dielectric constant, effective mass, electrical conductivity, electrical resistivity, energy bands, energy gap, energy levels, irradiation properties, lifetime, magnetoelastic properties, mobility, photo-electronic properties, photon, photoluminescence, piezoelectric properties, reflection coefficient and refractive index.
These data sheets present a compilation of a wide range of electronic properties for silicon. The energy band structure is thoroughly reviewed and includes effective mass, solubility, and diffusion with resulting values for energy levels and the dielectric constant. Electrical properties include mobility, resistivity, lifetime and piezoresistance. The various thermal and magnetic properties such as thermal expansion, thermal conductivity, phonon dispersion and the Seebeck, Nernst and Ettingshausen coefficients are reported. The g-factor, magnetic susceptibility and electron spin resonance are given. The optical properties, absorption, refractive index and reflectivity are reported as well as both photon and electron emission, including work function data. Each property is compiled over the widest possible range of parameters, including bulk and film samples. A data table containing a wide range of mechanical, physical and thermal properties is included, as well as a summary of crystal structure and phase transitions. There is also a general discussion of silicon technology.

The purpose of this symposium was to provide an introductory survey of those properties involved in the science and technology of materials. In the first chapter, Dr. C. O. Muhlhause describes the radiation fields present in reactors and the features of these fields most important to the development of nuclear materials, touching upon the principal physical effects induced in materials by the radiation. A. B. Liddiard, in the second chapter, discusses the solid state physics of radiation damage processes, with examples drawn partly from ceramics. The effects these processes have on physical properties are then described by P. Levy. Turning to chemical problems, R. J. Thorn and G. H. Windlow discuss the thermodynamics of non-stoichiometry, a particularly significant problem encountered with nuclear fuels. Finally, in the last two chapters, D. M. Read and J. H. Handwerk, and M. N. Morgan and D. Freshley take up the problems encountered in the development of nuclear fuels on the one hand, and structural and moderator materials on the other, thus displaying the applications of the basic ideas presented in the first four chapters.

These three volumes are collections of mineralogical data, the latest revision of a collection dating from 1837; the last edition was completed in 1892. The data were taken from earlier editions of the work, articles in mineralogical, chemical and physical journals, and independent books. Volume I (6500281) contains an introduction to the three volumes and data on native elements and sulfide, sulfosalts and oxide minerals; Volume II (6500282) contains data on halides, nitrates, borates, sulfates, phosphates, arsenates, tungstates, and molybdates. Volume III (6500283) is by C. Frondel only. It contains information on, and a discussion of, silica materials. The mineral data sections are arranged by the elemental composition of the minerals. The following properties are listed for each material: crystal structure and class, space group, axial ratios and angles, structure of unit cell, crystal habits, cleavage, hardness, specific gravity and color. Also, chemical analyses of specimens are presented, and data are provided on natural occurrence, artificial preparation of the mineral and the origin of its name.
the free ion are tabulated. These are applied to the
determination of the number and type of levels
arising from a free ion level with J ≥ 8. The
results of this analysis, together with the
selection rules for electric dipole, magnetic
dipole, and electric quadrupole transitions.
Calculation of the perturbation matrix elements by
the perturbation series and Racah coefficients is
discussed. Examples of the application of these
several techniques to specific problems are given.

Rand, N. H.; et al.
PLUTONIUM: PHYSICO-CHEMICAL PROPERTIES OF ITS
COMPOUNDS AND ALLOYS
Atomic Energy Review, 4:1-112
June 1966
This issue covers plutonium and its compounds and includes a critical evaluation of the data on thermodynamic properties, densities, crystallographic structures, compounds with non-metals, intermetallic phases, phase diagrams, equilibrium diagrams and diffusion rates in the condensed states. This assessment includes all the data published up to the end of 1964.

THE REACTOR HANDBOOK, VOLUME 5, SECTION 1: GENERAL
PROPERTIES OF MATERIALS
AEC-3647, Atomic Energy Commission,
Washington, D. C.
February 1955
610p
Available from Superintendent of Documents, GPO
This Handbook is a comprehensive and critical compilation of nuclear engineering data issued by the Atomic Energy Commission's Technical Information Service for reference use by scientists and engineers engaged in AEC reactor projects. The material presented represents the efforts of specialists in the various areas of reactor science and technology, and summarizes the accomplishments of the Commission's nuclear reactor program to date. Important nuclear reactor data are given in graphical and tabular form for the following metals and alloys: aluminum and its alloys; beryllium and its alloys; beryllia; beryllium carbide; bismuth; carbides; cements and concretes; glass; hydrides; lithium and its alloys; magnesium, molybdenum, nickel, plutonium, thorium, titanium, uranium, vanadium, zirconium, and their alloys; cobalt-base alloys; rare earths; silicon carbide; stainless steels; tungsten; and high-cross-section materials.

Reed, R. P.; Nokesell, R. P.
LOW TEMPERATURE MECHANICAL PROPERTIES OF COPPER AND
SELECTED COPPER ALLOYS, A COMPILATION FROM THE
LITERATURE
NBS Monograph 101, National Bureau of Standards,
Washington, D. C.
December 1967
161p
Available from the Superintendent of Documents, GPO
This compilation presents the mechanical properties of copper at temperatures up to 500K. The specific properties compiled are yield and tensile strength, elongation, impact energy, creep rate, reduction of area, hardness, fatigue behavior and moduli of elasticity and rigidity. The compilation is divided into four parts. The first section is intended for quick reference use for those who are interested in average values. The second section includes data from most of the investigators who have published results on the mechanical properties of copper and its alloys. The third section is composed of tables classifying the investigations which were not included in section two. These usually involve investigations in which data were obtained only at one temperature, such as room temperature. The fourth section lists, in alphabetical order, all references used.

Reiss, H.
PROGRESS IN SOLID STATE CHEMISTRY VOLUME 3
Oxford, New York: Pergamon Press
1967
510p
This third volume of Progress in Solid State Chemistry discusses the scientific understanding of the promising new techniques that have evolved for the preparation of solids in unusual states exemplified by single crystal epitaxial layers, solids at high pressure, and solids which are so disordered that they are essentially amorphous. The chapters include discussions on silicon heteroepitaxy on oxides by chemical vapor deposition; physical and chemical properties of semiconductor surfaces; solid state electrochemistry; lattice defects, ionic conductivity, and valence change of rare-earth impurities in alkaline-earth halides; non-stoichiometry in binary semiconductor compounds; lattice energies and related topics; phase relations and structures of solids at high pressures; metastable phases obtained by rapid quenching from the liquid state; and glass transitions in polymers. An author index, subject index, tables of data and references are included.

Cesium, S. J.; Ketchen, E. E.
CERIUM-90 DATA SHEETS
ORNL-4186, Oak Ridge National Laboratory, Oak Ridg e, Tennessee
December 1967
27p
Available from CFSTI
Data on Cs-137 are tabulated. Properties of the fuel forms CsCl, CsSO4, and cesium borosilicate glass are given. Twenty-seven references are included.

Cesium, S. J.; Ketchen, E. E.
CERIUM DATA SHEETS
ORNL-4187, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
32p
Available from CFSTI
Data on the fuel forms of Cm-244 are presented. Properties of Cm-244 metal, Cm02, Cm02S, CmF, Cm-242, 242Cm02 cermet are tabulated. Forty-nine references are given.

Cesium, S. J.; Ketchen, E. E.
STORIUM-90 DATA SHEETS
ORNL-4188, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
45p
Available from CFSTI
Data are given on composition, specific power, radiation, critical mass, container compatibility, thermal, physical properties, mechanical properties, chemical properties, biological tolerance, and shielding for strontium-90 metal, titanate (SrTiO3), oxide (SrO), fluoride (SrF2), and orthotitanate (SrTiO4). The data are indexed to 53 references which are also included.
This is an extensive revision and reworking of the second edition of the Space Materials Handbook. The Handbook is comprised of entirely new subject matter coverage and new materials data. All of the most significant material, phenomena, properties and principles covered in the original Handbook are presented and expanded in this revised and updated version. However, treatment of theoretical aspects has been condensed in order that more emphasis could be placed on the extensive new materials knowledge and data obtained from the design and successful launching of a wide variety of space systems. The Handbook is organized into four parts, namely: space environment, effect of space environment on materials, materials in space and biological interaction with spacecraft materials. Information on mechanical, physical and chemical properties and characteristics is given for a wide variety of metals, metallic and nonmetallic materials. The effects of natural and induced environments on materials are appraised. Materials categories include coverage of thermal control materials, optical materials, adhesives, organic structural materials, inorganic structural materials, electronic components and materials, materials for sealing applications and lubrication materials. In addition, a comprehensive multiple index is incorporated which gives ready access to information on specific subject areas with regard to their locations within the Handbook.

This is a noncritical compilation of data on superconductive materials that has been extracted from a portion of the literature published up to early 1968. The properties concerned are composition, critical magnetic field, crystallographic data, and lowest temperature tested for superconductivity. The compilation also includes a bibliography, general review articles and a special tabulation of high magnetic field superconductors. This compilation supersedes NBS Technical Note 408 and has the same title.

This monograph provides current information of the production, properties, and uses of high-purity and commercial forms of nickel and its important ferrous and nonferrous alloys. It is a revision of National Bureau of Standards Circular 592, issued in 1958. It gives 1180 references as well as tables and graphs of data.
These two reports present the results of a state-of-the-art survey covering tungsten, molybdenum and 10 of their alloys. All data are given in tabular and graphical form covering some of the more important physical, mechanical and metallurgical properties for each material. For both reports, the physical properties included are melting point, density, thermal expansion, thermal conductivity and electrical resistivity. The mechanical properties included are tensile properties (ultimate tensile strength, yield tensile strength, elongation, reduction in area, modulus of elasticity) and their temperature dependence, notched tensile properties, creep and stress-rupture properties, and other selected mechanical properties. Metallurgical properties include: fabricability, transition temperature, weldability, stress-relief temperature and recrystallization temperature. The ten alloy systems considered in the tungsten properties report include "doped" tungsten and various alloys with niobium, thorium dioxide, tantalum carbide, molybdenum and rhenium. The nine alloys considered in the molybdenum properties report include alloys with niobium, zirconium, titanium, tungsten and carbon. References are given at the conclusion of each materials section.


Available from CFSTI

This report is a compilation of the best available data on the physical properties of some of the plutonium compounds which are, or may be, of interest as fuel materials. These are the dioxide, monocarbide, monosulfide, mononitride, monoxide, mononitride, the dioxide, uranium-plutonium dioxide, uranium-plutonium monocarbide, uranium-plutonium mononitride, and the plutonium silicides.


Data on the properties of more than 520 different materials, such as carbides, borides, nitrides, sulfides, phosphides, silicides, oxides, as well as mixed oxides and mixed carbides are presented. The data are taken from 698 references, with the large majority from the U.S. Government report literature. The data concern general, chemical, electrical, mechanical, nuclear, optical, structural, and thermal properties. The references are listed by number as they appear in the tables, as well as alphabetically by name of author. The extensive table of contents on pages vii to xx makes it easy to find the various materials in the tables.


The data contained in Data on Chemicals for Ceramic Use are a revision of Bulletin 197, which was taken from the Report of the Committee on Chemical Data for Ceramists submitted to the Division of Chemistry and Chemical Technology of the National Research Council, 1942. Elements and compounds are arranged alphabetically according to names. All available data as of 1949 on density, melting point, transition point, boiling point, sublimation point, decomposition temperature, refractive index, crystal form, colors, formulas and molecular weights is listed. This revised edition brings the report up to June 1949.

Simonds, H. R.; Church, J. H. *8.373-4.373* A CONCISE GUIDE TO PLASTICS, SECOND EDITION New York: Reinhold Publishing Corporation 1965 404p

This second edition brings up to date the text of the first edition of A Concise Guide to Plastics and encompasses a much larger field due to the expansion of the plastics industry itself. An objective of this book was to answer the questions asked at the 1961 Plastics Exhibit. The subjects discussed include basic chemistry of plastics, manufacturing techniques, stereospecific catalysts, ablative action, borderline materials, forms of plastics, resin manufacture, compounding, processing, applications, production and prices, selection factors, and the business of plastics. One chapter is devoted to plastic manufacturers' statements and contains information about the company's materials, trade names and sales for fifty-two producing firms. Over fifty pages of tables of the United States Plastics Trade Names are included, as well as numerous graphs and tables of data concerning plastic properties.
These two books contain data on the X-ray powder diffraction patterns for many compounds. The inorganic and organic volumes were prepared by photographing file cards, each of which contains data on one compound. The data given for each compound includes the interplanar spacings corresponding to the three strongest lines in the spectrum and the largest interplanar spacing found for the specimen; the intensities of these lines; wavelength, source and filtered of the X-rays used; crystallographic system, lattice parameters and interaxial angles; indices of refraction, density, melting point and color of the specimen; chemical and structural formulas; interplanar spacing, intensities and Miller indices for all lines found; and the reference to the literature from which the data were taken.

**References:**


These volumes represent a continuing NBS-ASTM project for improving a file of X-ray diffraction patterns. Reports are made on substances for which powder patterns are desired and added to the file.

The ASTM patterns are tabulated with additional patterns from the literature and one prepared at the NBS. Miller indices are derived from the calculation of spacings by desk calculator or the electronic computer SEAC. Interplanar spacings in angstroms (except where otherwise noted) and relative intensities from 0 to 100 are tabulated. For the NBS pattern the three strongest lines are given, as well as the lattice constants and the computed density. The index of refraction of the sample is noted if it could be determined. Crystal-structure data from the literature are noted. There fifty-four substances covered in Volume I; Volume II includes thirty substances; Volume III includes thirty-four substances; Volume IV includes forty-five substances; Volume V includes forty substances; Volume VI includes sixty-one substances; Volume IX includes forty-three substances; and Volume X includes forty substances. A Cumulative Index to Volumes I-X is located at the end of Volume X.
The tensile and impact properties of structural materials were experimentally determined at temperatures from 20 to 300°K. Tensile properties of a few materials were also determined at 4°K. The materials included forty-two commercial alloys of iron, aluminum, titanium, copper, nickel, and cobalt, and two metal-bonded carbides. The properties experimentally determined were the yield strength, tensile strength, elongation, and reduction of area, the stress versus strain curve, and the impact energy. The test equipment and procedures are described. The individual data are presented in tables, and the average results are displayed in graphs.

These data sheets present a compilation of the electronic, thermal and optical properties of copper over the widest possible range of parameters from references obtained in a thorough literature search. These properties are correlated with each other and with the theoretical and experimental description of the Fermi surface of copper. An extensive evaluation of the electrical resistivity of pure copper is presented. Data are given on magnetoresistance, Hall effect, superconductivity, thermal conductivity, Lorenz number, thermoelectric power, specific heat, optical properties, and electron emission. Representative data on many copper alloys and the anomalies observed in the properties of dilute copper-transition metal alloys are considered. A detailed index to the data is presented.

This second edition includes X-ray emission line wavelengths in angstrom units for the range 0.25 to 100 angstroms. The table is especially designed for use with curved crystal spectrometers that are geared to read directly in wavelength. The table is presented in two sections. Section one lists all of the first order lines including satellite and low intensity diagram lines for each element (boron and above), arranged in order of increasing atomic number with lines sorted on the basis of decreasing wavelength within each element. Section two lists all lines sorted on the basis of increasing wavelength. Odometer settings for commonly interchanged crystals have been computed and listed for each line in both sections. The two sections are separated by a periodic table of the elements which lists the most intense lines for each element.
the III-V compounds. Contributions from ten to fifteen scientists are included in each volume. The books are designed for reference work as well as texts on the graduate level. Volume 1 reviews key properties of the III-V compounds, with special emphasis on band structure, magnetic field phenomena, and plasma effects. In volume 2, the emphasis is on physical properties, phenomena, magnetic resonances, and photoelectric effects, as well as radiative recombination and stimulated emission. Volume 3 is concerned with optical properties, including lattice effects, intrinsic absorption, free carrier phenomena and photovoltaic effects. Volume 4 includes thermodynamic properties, phase diagrams, diffusion, hardness, and phenomena in solid solutions. Volume 5 is concerned with the effects of strong electromagnetic fields, hydrostatic pressure, nuclear irradiation and nonuniformity of impurity distribution on the electrical and other properties of III-V compounds.

Each point has a key number which identifies the compound.

Winchell, A. N.; Winchell, H.
ELEMENTS OF OPTICAL MINERALOGY: AN INTRODUCTION TO MICROSCOPIC PETROGRAPHY. PART 2, DESCRIPTIONS OF MINERALS, FOURTH EDITION
New York: John Wiley & Sons, Inc.
1961
551p

This book is a compendium of data on the physical and crystalline properties of minerals taken from literature published through 1949. The authors have presented data on the properties of these materials as a function of their composition in elements and simple molecules. The data included for each entry are: mineral name; chemical formula with known analytical variations; crystal structure symmetry and space group; lattice constants and axial angles; hardness; specific gravity; melting point; indices of refraction along principal axes; and optical angles; specific refraction; and tables of these properties as a function of composition are provided for materials of variable composition. Data are provided for native element substances and for halides, sulfides, oxides, carbonates, borates, sulfates, phosphates and silicates.

Winchell, A. N.; Winchell, H.
THE MICROSCOPICAL CHARACTERS OF ARTIFICIAL INORGANIC SOLID SUBSTANCES; OPTICAL PROPERTIES OF ARTIFICIAL MINERALS, THIRD EDITION
1964
439p

This book contains descriptions of inorganic solid substances whose properties are known and a set of determinative tables and charts for identifying an unknown compound from its optical properties. The following data are given for each substance: chemical formula and name; dimensions and axial angles of the unit cell; data on crystal habits, cleavages; and axial relations; specific gravity; optical properties, including optic orientation, principal refractive indices, optic sign, optic axial angle, dispersion, color, and absorption. The spacings for the substance's X-ray powder diffraction pattern are also given. The determinative tables are presented as polar plots of optic angle vs. birefringence with each plot being photographed on a polar plot of optic variation. Each compound is represented as a point on such a plot, and a table accompanying each plot gives a page number for the text where the compound can be found. The following classes of compounds are included: elements, carbonates, nitrides, sulfides, selenides, tellurides, sulfoalts, halides, hydrates, amides, cyanides and cyanates, oxides, carbonates, nitrites, halides, borates, sulfates, nolybdates, tungstates, phosphates and silicates.

Winchell, A. N.
OPTICAL PROPERTIES OF MINERALS: A DETERMINATIVE TABLE
1965
91p

This compilation presents data useful for the optical identification of minerals. The data include the refractive indices, birefringence, and optic axial angle for over 1000 minerals. The data are arranged in alphabetical order of the compound name. The values of the optic angle against birefringence for materials falling within a narrow range of refractive indices. Each mineral is represented by a point on such a graph. Also listed for each material are the strongest
A critical survey has been made of published values for sixteen of the principal physical constants of rubber in the fields of mechanics, heat, optics and electricity, namely: dimensions of unit cell, density, thermal conductivity, specific heat, heat of fusion of crystalline rubber, heat of combustion, volume compressibility, Poisson's ratio, velocity of sound, refractive index and dispersion, stress-optical coefficient, dielectric constant, power factor, and conductivity. The value for each constant which seems to be the most reliable has been indicated, and, when necessary, corrected to the standard conditions of normal atmospheric pressure and a temperature of 25°C. Temperature coefficients are given when possible. The values are given for rubber in four different forms: the purified hydrocarbon, commercial raw rubber, soft vulcanised rubber containing 24 combined sulfur, and hard rubber containing 52 combined sulfur. Relationships between expansivity and change of density with temperature, expansivity and change of refractive index with temperature, and refractive index and dielectric constant are discussed.

This report is a compilation of data on thermal radiative properties. It also includes a brief discussion of the basic fundamentals of thermal radiation and of the methods of measuring these properties. Much of the information has previously been distributed in NHC memorandum; however, it is consolidated in this report for the benefit of those with a broad interest in radiant heat transfer. Thermal radiative data are included for the following materials: titanium and its alloys; stainless steel, iron, nickel, cobalt-base superalloys; the refractory metals (chromium, columbium, molybdenum, tantalum, and tungsten) and their alloys; coated materials for elevated-temperature service; and ceramics and graphite. Forty-one references are included.

These volumes provide collections of data taken from the literature on the structure of crystals. Information is provided in the form of text, tables and illustrations. Data provided in the tables for each crystal include: crystal type and class, and its lattice constants. For each crystal type, data on symmetry and positions of atoms within the lattice are provided in the text section, and illustrations of the typical compounds are given. The volumes are divided into chapters according to the chemical type of the compounds. Volume I contains elements and compounds of the type RX and RXR. Volume II contains complex binary compounds RMX, and structures of the type R(MX)2, R(MX)3, and Rn(MX)m. Volume III contains hydrates, ammoniates, silicates and miscellaneous inorganic compounds plus an inorganic formula index and a mineralogical name index. Volume IV contains structures of aliphatic compounds, while Volume V contains structures of benzenoid derivatives and alkylic and heterooid containing carbon hydrates, and an index to organic compounds.

The book is divided into four chapters: preface and introduction; structures of the elements; structures of the compounds RX; structures of the compounds RX2. A bibliography is given at the end of each chapter. A name index and formula index are provided at the end of the book.

This book is divided into three chapters: structures of complex binary compounds RX; structures of the compounds RXn and compounds of the type R(MX)2. A bibliography is given at the end of each chapter. A name index and formula index are provided at the end of the book. This attempt to provide a unified statement of the atomic positions in all analyzed crystals progresses, improvements become possible. The principal one introduced with this volume is a more reliable bibliography.

This third volume contains three chapters giving crystal structure for inorganic compounds of the type R(MX)2, type R(MX)3, and hydrates and ammoniates. A bibliography is given at the end of each chapter. The book also includes name and formula indexes.

About 550 references are contained in this bibliography which includes, it is believed, a complete coverage of the soft X-ray literature since 1950 and through 1960. Some references to earlier work are given, but the general review references, listed separately, give an adequate coverage of the earlier work. The emphasis is on the application of soft X-ray spectroscopy in the study of valence band electronic states in metals and alloys, and therefore, the spectral region of 25 to 100 angstroms involving ruled glass grating spectrometers is of principal interest. However, a wealth of data have been gathered, primarily by the Russians, on valence electronic states by means of high energy transitions where crystal spectrometers are satisfactory. These references and any X-ray work leading to the distribution of valence
Electronic states are included regardless of the transitions employed. In addition to soft X-ray data, references on all pertinent aspects of the apparatus and experimental problems are included. Also listed separately are references of value in corroborating soft X-ray data with other results, such as energy band calculations. Subject, author, X-ray band, material, and other indices are included.


The bibliography is presented in four principal parts: Part I is a general bibliography of 4500 entries arranged alphabetically by authors, but not including the writings of P. W. Bridgman. Part II is an index of the material of Part I, arranged by subject. Part III is a complete bibliography of the 310 writings of P. W. Bridgman. Part IV is an index of the material in Part III, arranged by subject.
Chemical Kinetics
Amis, E. S.
SOLVENT EFFECTS ON REACTION RATES AND MECHANISMS
New York: Academic Press
1966
228p

Theoretical explanations are given for the solvent effects on reactions between various charged systems of reactants. These include ion-ion, ion-dipolar molecule, dipolar molecule-dipolar molecule, exchange, and some charge type reactions which do not fit in the above classes. Hypothetical solvent effects as well as experimentally observed effects are explained on a theoretical basis.

Bates, P. R. (Editor)
ATOMIC AND MOLECULAR PROCESSES
New York: Academic Press
1962
904p

This compilation, which is designed primarily as a reference book for chemists and physicists concerned with radiative and collisional processes involving atoms or molecules. It provides surveys covering the following topics: forbidden and allowed transitions; bands, photoionization, photodetachment; recombination, attachment; elastic and inelastic scattering of electrons, energy loss by slow electrons; collision broadening of spectral features; electron transfer between atomic systems including range, energy loss, excitation, ionization, detachment, charge transfer, elastic scattering, mobility, diffusion, relaxation in gases, and chemical reactions. A chapter is devoted to the use of high-temperature shock waves and accounts are given of the other main experimental methods. The relevant theoretical work is also described, detailed mathematics being avoided as far as possible.

Baulch, D. L.; et al.
HIGH TEMPERATURE RATE DATA, CRITICAL EVALUATION OF RATE DATA FOR HOMOGENEOUS, GAS-PHASE REACTIONS OF INTEREST IN HIGH-TEMPERATURE SYSTEMS
Reprinted Nos. 1-5, 7, 21, 3, Department of Chemical Physics, The University of Leeds, England
May 1968, December 1968, April 1969
25p, 34p, 54p

These reports are a series of critical evaluations on gas-phase reaction data. They contain discussions of the values listed and include bibliographies. Report No. 1 treats the following reactions:

\[ \text{CO} + \text{OH} = \text{CO}_2 + \text{H} \]
\[ \text{CO} + \text{OD} = \text{CO}_2 + \text{D} \]
\[ \text{CO} + \text{O} = \text{CO}_2 + \text{hnu} \]
\[ \text{CO} + \text{O} = \text{CO}_2 \]
\[ \text{CO} + \text{O} + \text{M} = \text{CO}_2 + \text{M} \]
\[ \text{CO}_2 + \text{O} + \text{CO}_2 \]

Report No. 2 treats the following reactions:

\[ \text{H}_2 + \text{O} + \text{OH} + \text{H} \]
\[ \text{H}_2 + \text{OH} = \text{H}_2 + \text{O} \]
\[ \text{H}_2 + \text{O}_2 + \text{OH} + \text{H} \]
\[ \text{H}_2 + \text{O}_2 + \text{OH} + \text{H} \]
\[ \text{O} + \text{OH} + \text{H}_2 + \text{O} \]
\[ \text{O} + \text{OH} + \text{H}_2 + \text{O} \]

Report No. 3 treats the following reactions:

\[ \text{O}_2 + \text{H} = \text{O} + \text{OH} \]
\[ \text{O} + \text{OH} = \text{O}_2 + \text{H} \]
\[ \text{O}_2 + \text{H} + \text{M} = \text{H}_2 + \text{O}_2 + \text{M} \]
\[ \text{H}_2 + \text{H} + \text{M} = \text{H}_2 + \text{O}_2 + \text{H} + \text{M} \]
\[ \text{H}_2 + \text{H} + \text{M} = \text{H}_2 + \text{O}_2 + \text{H} + \text{M} \]
\[ \text{H}_2 + \text{H} + \text{M} = \text{H}_2 + \text{O}_2 + \text{H} + \text{M} \]

Berlman, I. B.
HANDBOOK OF FLUORESCENCE SPECTRA OF AROMATIC MOLECULES
New York: Academic Press
1965
258p

Spectra of approximately 100 aromatic molecules, varying in size, shape, and structure, are assembled in the handbook. The list of molecules begins with the simple aromatic molecule benzene and includes progressively more complicated systems. Although a special effort has been made to include practically all of the currently popular organic scintillators, the choice of the other molecules has been rather arbitrary. The luminescence characteristics of each aromatic molecule have been measured in a systematic fashion. Fluorescence and absorption spectra are plotted for each molecule and additional data related to the fluorescence process are given in each graph. The additional data include the fluorescence decay time, the fluorescence quantum yield, the natural lifetime as computed from the absorption spectrum, the Stokes loss, the wavelength of the center of gravity of the fluorescence spectrum, and the average wavelength of the fluorescence spectrum. Some supplementary material as topical bibliographies, tables to convert wavelength to wave number and energy, a table of values of the index of refraction of cyclohexane as a function of wave number, and a table of values of oscillator strengths have been placed in the Appendix.

Berry, R. S.
SMALL FREE NEGATIVE IONS
Chemical Review, 69:533-542
August 1969

This article surveys the restricted subject of properties of isolated atomic and small molecular negative ions. Included in the discussion are electron affinities, electronic states, optical properties, and the experimental and theoretical methods for studying these properties. Excluded from the discussion is the vast group of problems associated with large polyatomic negative ions and with most of the rich subject of collision processes involving negative ions, such as detachment and transfer of electrons in collisions.

BERLINER, L. (Editor)
BIBLIOGRAPHY ON THE RADIATION CHEMISTRY OF AQUEOUS SOLUTIONS - I. AMINO ACIDS 1952-1968
COO-38-641, Radiation Chemistry Data Center, Radiation Laboratory, University of Notre Dame, Notre Dame, Indiana
November 1968
16p
Available from CFSTI

This bibliography, covering the literature from 1952 to 1968, contains radiation data for amino acids. Abstract references are included for foreign language articles and technical reports. The papers are listed alphabetically by year with each year's papers listed alphabetically by first author.

BERLINER, L. (Editor)
BIBLIOGRAPHY ON THE RADIATION CHEMISTRY OF AQUEOUS SOLUTIONS - II. CARBOHYDRATES 1952-1968
COO-38-642, Radiation Chemistry Data Center, Radiation Laboratory, University of Notre Dame, Notre Dame, Indiana
December 1968
11p
Available from CFSTI

This bibliography covering the literature from 1952 to 1968 contains radiation data for carbohydrates. Abstract references are included for most foreign language articles and technical reports. The
papers are listed by year, with each year's papers listed alphabetically by first author. At the end of each reference the Radiation Chemistry Data Center serial number is given.

BIBLIOGRAPHY ON THE RADIATION CHEMISTRY OF ALCOHOLS AND PHENOLS
C00-38-643, Radiation Chemistry Data Center, Radiation Laboratory, University of Notre Dame, Notre Dame, Indiana November 1968 34p Available from CFSTI

This bibliography covering the literature from 1953 to 1968 contains radiation data for alcohols and phenols. Abstract references are included for foreign language articles and technical reports. Investigations of aqueous solutions of alcohols and phenols are not included; polyhydron compounds (glycols) are also not included. The papers are listed under the categories by serial number which is approximately chronological. If an article is pertinent to more than one category it is listed under each. The compound index includes the alcohols or phenols and additives, and refers to the serial number of the paper.

BLANK, C. A. (Editor)
DASA REACTION RATE HANDBOOK
Santa Barbara, California: DASA Information and Analysis Center, TEMPO October 1967 410p

Chemical rate processes are of extreme importance in the deionization and recovery of the highly ionized atmosphere created by a nuclear burst. The Handbook presents what it considers to be the best available chemical kinetic data and related unclassified information, required to calculate the kinetics of atmosphere deionization. The Handbook is divided into four major sections, each comprising several chapters of related information. The first of these sections deals with general background materials. The conditions of temperature, density and composition which are needed to determine the importance of specific processes are discussed for the normal and disturbed atmosphere. Basic data, energy levels and equilibrium constants are given and the general types of processes which need to be considered are discussed. In the second section, various aspects of the different methods employed to obtain data on the processes of importance are discussed. The nature and scope of such data are considered in the third section of this Handbook, where the state of knowledge of various types of reactions is examined and evaluated. The fourth section contains chapters dealing with the proper utilization, status, uncertainties and central problem areas associated with the determination of reaction rates as well as a detailed summary of values recommended for use in atmospheric deionization calculations.

SORNER, M. H.
A REVIEW OF RATE CONSTANTS OF SELECTED REACTIONS OF INTEREST IN RE-ENTRY FLOW FIELDS IN THE ATMOSPHERE

The major reactions normally encountered in flow field re-entry calculations for the oxygen-nitrogen system are reviewed and a rate constant value for each reaction is recommended. Collisional ionization processes, charge-exchange reactions and attachment-detachment reactions are included. Rate constant coefficients to fit the equation

\[ k = aT^b e^{-c/T} \]

are reported in tabular form for the selected rate constant. The data are also graphed as log k vs T(K). In all, twenty forward and reverse reactions are reviewed. Additionally, a large number of rate constant values for reactions which also effect flow field calculations are given.

COHN, C. E.; GOLDEN, G. H.
A TABLE OF THERMODYNAMIC PROPERTIES OF HYDROGEN FOR TEMPERATURES FROM 100 TO 5000K AND PRESSURES FROM 1 TO 50 ATMOSPHERES
ANL-6673, Argonne National Laboratory, Argonne, Illinois January 1963 267p Available from CFSTI

This report presents a table of thermodynamic properties of gaseous hydrogen covering the temperature range from 100 to 30000K and the pressure range from 1 to 50 atmospheres. The calculations have been done by means of a computer routine. The table has entries for temperature at 10K increments, for pressure at 1 atmosphere and at 5 atmosphere increments from 5 through 50 atmospheres, and for 25%, 50%, 75%, and 100% of the gas initially in the para state. The first of these latter values corresponds to normal hydrogen, whereas the remaining values apply to vapor from liquid which has been stored for varying lengths of time. For each entry, the table gives the density in gm/liter, the enthalpy in joules/gm, the entropy in joules/(gm)(K) and the fraction of the gas molecules which are dissociated.

COOK, C. J.; LORENTS, D. C.
ELECTRON COLLISION FREQUENCIES AND SCATTERING CROSS SECTIONS IN THE IONOSPHERE
A Final Technical Report No. 6, Stanford Research Institute, Menlo Park, California August 1961 54p Available from DDC AD 291 591

The propagation of electromagnetic waves through an ionized, gaseous medium can be described in terms of a set of macroscopic parameters that are a function of the physical properties of the medium, the permittivity, conductivity, and permeability. This report calculates the propagation parameters for the normal ionosphere above 50 km, and from these parameters calculates theoretical values for electron collision frequencies and scattering cross sections in the ionosphere.

COOPER, J. (Editor)
PROCEEDINGS OF WORKSHOP CONFERENCE ON THE LOWERING OF THE IONIZATION POTENTIAL AND RELATED PROBLEMS OF THE EQUILIBRIUM PLASMA
JILA Report No. 79, Joint Institute for Laboratory Astrophysics, Boulder, Colorado July 1966 225p

This workshop was sponsored by the Joint Institute for Laboratory Astrophysics. Its concern was statistical mechanics of plasma both from experimental and theoretical points of view. There were many recent development and advances in the treatment of such problems as the plasma partition function and the lowering of the ionization potential, although there were strong disagreements between different workers in this field. The proceedings were taped and should be considered as unpublished material. It was the object of this workshop to bring together active theorists and experimentalists in this field to resolve some of the differences. Included in the Appendix is a bibliography on Research on Lowering of the Ionization Potential.
These bibliographies consist of references to research involving temperatures above 1000°C, which were noted by the contributors during the above-month and three-month periods; suffice are intended primarily as current-awareness bibliographies, there is no cross-referencing or indexing. These issues contain about 700 references each, roughly grouped under fifteen subject headings.

Dragoo, A. L.

DIFFUSION RATES IN INORGANIC NUCLEAR MATERIALS

The tracer diffusion coefficient, the self-diffusion coefficient, and the interdiffusion coefficient are briefly described. Grain boundary and lattice (volume) diffusion are contrasted. The frequency factors (Q) and activation energies (Q) are tabulated for diffusion in the borides, carbides, and oxides of Be, Hf, Mo, Nb, Ta, Th, Ti, and Zr and for diffusion of C, N, and O in these metals. The purity of the solvent media, the preparation and properties of the samples, the method, the type of diffusion coefficient measured, and the temperature range are also specified.

Flynn, J. H.

RAPID DETERMINATION OF THE ORDER OF CHEMICAL REACTIONS FROM TIMED-RATIO TABLES

A table of ratios of reaction times corresponding to the time (t) and final time (t + 0) is presented. Use of this table allows the rapid calculation of the order of chemical reactions that are experimentally simple and involve only one subjective step. Methods are discussed for interpreting deviations from constant order resulting from errors in the initial times and concentrations, from errors in stoichiometry, and from the reaction following a more complicated rate expression.

Franklin, J. L.; et al.

IONIZATION POTENTIALS, APPEARANCE POTENTIALS, AND HEATS OF FORMATION OF GASEOUS POSITIVE IONS
NBS-NSR-26, National Bureau of Standards, Washington, D.C., June 1969

This is a compilation of ionization and appearance potentials of positive ions published from 1955 to June 1969. The compilation lists the ion formed, the parent species from which it was formed, the other products of the process, the threshold energy for the formation of this ion, and the method by which this data was obtained. Where feasible, the heat of formation at 298 K of the positive ion has been computed for each entry using auxiliary thermochemical data. From these computed values "best" values have been chosen.

Garvin, D. (Editor)

A COMPENDIUM OF EVALUATED AND ESTIMATED RATE COEFFICIENTS
July 1968
316p

This report is an annotated handbook on rates of elementary gas phase reactions. It is limited to rate evaluations. No attempt has been made to record all research papers pertinent to a particular reaction. There are two principal parts to the report: a compilation and a set of rate coefficients. The compilation merges the information presented in several recent journal articles and reports that are devoted to the evaluation or estimation of rate coefficients for elementary reactions. A single table, arranged by reaction, lists the various recommended rates in a common format. Notes to the table show what data sources were used by each evaluator, and, where possible, how his recommended value was determined. The person requiring a rate for a specific reaction may compare the recommended values, compare their data bases and also extract a bibliography of the pertinent work. Rate evaluations which are presented in full, stand independent of the main table. They are for reactions of some importance to combustion and atmospheric chemistry. They are presented in a form suited to the reaction-by-reaction evaluation of each data as opposed to that used in monographs. It is hoped that they will serve as models, albeit ones to be improved upon, for future rate data examinations.

Harllee, F. N.; Rosenstock, H. M.; Herron, J. T.

A BIBLIOGRAPHY ON ION-MOLECULE REACTIONS, JANUARY 1900 TO MARCH 1966
June 28, 1966
38p

This document is a bibliography on ion-molecule reactions of chemical interest. The bibliography is listed in a chronologic-alphabetic arrangement and includes 451 titles and references taken from the published literature since 1900. The coverage is limited to reactions between ions and molecules leading to changes in the atomic composition of either the projectile ion or target molecule. No attempt has been made to completely cover resonant charge exchange or elastic scattering.

Henley, E. J.; Johnson, E. R.

THE CHEMISTRY AND PHYSICS OF HIGH ENERGY REACTIONS
1969
475p

This is a text on radiation chemistry on the graduate level, discussing radiation effects on complex systems. The book includes discussions on general nomenclature; interaction of radiation with matter; radiation sources and dosimetry; chemical consequences of the absorption of high energy radiation; matter-dose data in radiation chemistry, radiation chemistry of gases, water, and of aqueous and solid systems; effect of radiation on liquid hydrocarbons and polymers; radiation-induced polymerizations; and industrial applications.
Tables of sin E and l-cos E with argument E = sin E, where parallel tables in the hyperbolic functions are given for the determination of position and velocity from the time, for rectilinear orbits. Direct interpolation is possible, without the aid of series expansions or successive approximations. The tables may also be used in connection with nearly rectilinear motion, that is, motion in ellipses and hyperbolas whose excentricities are nearly unity. The application of the tables is set forth extensively in the Introduction.

Hochstim, A. R. (Editor)
Berman, M.; et al.
BIBLIOGRAPHY OF CHEMICAL KINETICS AND COLLISION PROCESSES

This book contains about 20,000 entries, each specifying the study of a collision process, drawn from about 7000 selected articles. The entries cover the following: reaction rates of atoms, ions, and small molecules in the gas phase; inelastic scattering cross sections of atoms, ions, and small molecules in the low-energy range (E < 100 eV); low-energy elastic electron collision cross sections; lifetimes of excited states of various species; and reaction rates involving solid carbon. In the context of this book, "small molecules" means species with four atoms or less. Only occasionally are larger molecules included.

Horiuti, J.; Miyahara, K.
HYDROGENATION OF ETHYLENE ON METALLIC CATALYSTS

Available from Superintendent of Documents, GPO

Reaction rate data for the catalyzed hydrogenation of ethylene, primarily in the presence of unsupported metallic catalysts, are critically reviewed. Reaction mechanisms are discussed in detail, and a statistical mechanical treatment of the reaction is given, according to the generalized theory of reaction starting from the well-known procedure of Glastone, Laidler, and Eyring. Data for single-element catalysts and alloys are included and interpreted, as are data illustrating differences due to the physical form of the catalyst (film, foil, wire, powder, and some supported systems). Problems are discussed concerning reproducibility of experimental results over repeated runs, and as a function of catalyst pretreatment. The data are analyzed in 29 graphs and 29 tables, some of which are very extensive. The bibliography includes 141 references.

Johnston, Harold S.
GAS PHASE REACTION KINETICS OF NEUTRAL OXYGEN SPECIES

Available from Superintendent of Documents, GPO

The available data for reactions among neutral oxygen species, oxygen atoms, oxygen molecules and ozone, have been reviewed. Selected data have been reanalyzed and used to establish values for the rates of these reactions:

\[
\begin{align*}
0 & + 2 \cdot M + 0 + 0 + M \quad (M = 02, Ar) \\
0 & + 0 + M + 02 + M \quad (M = 02) \\
0 & + M + 0 + 02 + M \quad (M = 02, 03, He, N2, CO2) \\
0 & + 02 + M + 03 + M \quad (M = 02, 03, He, Ar, N2, CO2) \\
0 & + 03 + 02 + 02 \\
0 & + 02 + 0 + 02 \\
\end{align*}
\]

(isotopic exchange)

Johnston, H. S.
GAS PHASE REACTION RATE THEORY

This is a college text which reviews the theories of chemical kinetics based on molecular mechanics and chemical kinetics. The algebraic structure of simple two-atom collision rates and three-atom metathetical reactions is presented in detail. Considerable use is made of graphical presentation in terms of three-atom models. More complex methods are briefly outlined and reference is given to advanced publications. Several unpublished studies are included. Experimental data are compared with theory whenever possible. Only a few chemical reaction systems are discussed in detail. Tables of chemical kinetics data and a large number of references are included.

Jones, R. N.; et al.
COMPUTER PROGRAMS FOR ABSORPTION SPECTROPHOTOMETRY
NRC Bulletin No. 11, National Research Council of Canada, Ottawa, Canada 1968 159p

Nine "building block" computer programs for performing the basic numerical computations of absorption spectrophotometry are described and listed. They are written in FORTRAN IV for card input and output. The input and output formats are standardized to permit easy interfacing to produce more complex data processing systems. Though developed for infrared spectrophotometry these programs are adaptable for use in the visible and ultraviolet. The operations covered include scale change and scale interval change, location of band maxima and minima, smoothing, second derivative, slit function convolution and deconvolution, single band generation, profile analysis and asymmetry analysis of single bands from the truncated moments, and ordinate addition.

Kerr, J. A.
BOND DISSOCIATION ENERGIES BY KINETIC METHODS
Chemical Reviews, 66: 465-500 September 26, 1966

The main aim of this review is to present an assessment of bond dissociation energies, which have recently been determined for the first time. The review is centered on polyatomic molecules, and deals with modifications and expansions of the methods which have been developed over the past few years. The bond dissociation energy or bond strength of a chemical bond A-B is usually taken to mean the enthalpy change of the reaction A-B = A+B in the ideal gas state referred to 298K. Experimental methods are given for pyrolytic reactions, metathetical reactions, shock-tube reactions, and estimated errors. Data on bond energies are given for hydrocarbon, organic nitrogen-oxygen-, sulfur- and halogen-containing compounds, and metal alkyls. Data are given for heats of formation of atoms and radicals, bond dissociation energies and best available values of bond dissociation energies.
A comprehensive compilation of low energy electron collision cross-section data is presented. This is the first of three parts and is limited to experimental measurements, including data for all atomic species and for those molecules which are important in aeronomy, astrophysics and plasma physics. Literature data through September 1968 are included. The compilation is divided into three main categories: ionization, including all processes which lead to electron ejection from atomic or molecular targets; dissociation, including processes which lead to ionized fragments; and vibrational and rotational excitation. The data are presented in order: atoms, diatomic molecules, triatomic molecules, etc. The atoms are ordered by the Z number, and the molecules are ordered according to the atom with the largest Z in the molecule.
The number of ab initio molecular electronic calculations has increased dramatically in the last few years. Both the practitioners and other interested students of the results of the calculations have found it increasingly difficult to determine the present status of these calculations. This compendium references the work from 1960 to the present and abstracts from the mass of data the best values for several observable properties in individual electronic states: ionization energy, electron affinity, spectroscopic constants, electric moments, field gradients, polarizabilities, and magnetic constants. In order to provide an insight into molecular electronic structure, tables of orbital energies are also included. These tables are meant to direct attention to the successes and failures of the calculations by comparing a large percentage of the best results in a reasonably compact form. Its usefulness will be limited in time by rapid advance in the field.

#6906001 *3.153-5.153
Laborie, P.; Rocard, J.-M.; Rees, J. A.
ELECTRONIC CROSS-SECTIONS AND MACROSCOPIC COEFFICIENTS: 1. HYDROGEN AND RARE GASES
Paris: Dunod 1968 199p In French and English

This catalogue contains some of the principal experimental data concerning the interaction of electrons with atoms and molecules. It is a non-critical selection of data from the most accurate and most recent measurements available. The cross-sections are all expressed in cm² per atom or molecule, and given as a function of the electron energy expressed in electron volts. Curves are reproduced where the original article gives data in this manner. Tables of data are given where available. Atomic or molecular energy levels and macroscopic coefficients are included. The book includes data for hydrogen, helium, neon, argon, krypton, and xenon.

#6500526 *0.1151-3.151-5.151-8.151
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME 1: ATOMIC AND MOLECULAR PHYSICS, PART 2: MOLECULES I (NUCLEAR STRUCTURE)
Eucken, A.; Hellwege, K.-H. (Editors) Berlin, New York: Springer-Verlag 1951 571p In German

This section of the Landolt-Bornstein tables contains atomic physical data of molecules which describes the structure and dynamics of their nuclear framework. The following properties have been tabulated: atomic distances and structures, valence energies of chemical bonds, dissociation energies of chemical bonds, and molecular vibrations and rotations.

#6630060 *0.1151-5.151-8.151
LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME II: PROPERTIES OF MATTER IN ITS AGGREGATED STATES
PART 5b: TRANSPORT PHENOMENA II, KINETIC HOMOGENEOUS GAS EQUILIBRIA
Schaefer, K. (Editor)
Berlin, New York: Springer-Verlag 1968 397p In German

Part 5b of Volume II contains some kinetic properties of the elements and selected compounds. The first section lists diffusion constants for elements and ions in metals, salts, and alloys, both liquid and solid. A short table of values for diffusion constants of some gases in selected metals is included. The second section of tables deals with thermal conductivity data for gases, liquids and solids. The constants are reported as a function of temperature, pressure and composition. The next series of tables deals with thermal diffusion constants for binary and ternary gas mixtures. Constants are also reported in liquid mixtures and solutions (Ludwig-Soret effect). The next section lists a series of kinetic rate constants for reaction in gas phase and includes: dissociation, atom-radical and ion-molecule reactions. Another series of tables lists rate constants and equilibrium constants for reactions in solid and mixed phase. Several addenda to the main sections are appended.
Available theoretical and experimental data on the reactions that control the electron density in missile wakes and in the environment surrounding nuclear bursts have been reviewed. Calculations of cross-sections (reaction rates) for specific reactions of importance in these problem areas have been made. The aim of the work here is also to develop general computational techniques designed to improve the accuracy and reliability of calculations.

#6002032
Motaia, R. A.
COMBUSTION KINETICS OF TETRAFLUOROETHYLENE
Final Technical Report, Fluid Dynamics Laboratory, Department of Mechanical Engineering, University of Michigan, Ann Arbor, Michigan
July 1968
41p
The pyrosis of C2F4, C3F6, and CF2O have been studied in the temperature and pressure ranges 300-455 °C, 25-760 torr; 550-675 °C, 50-410 torr; and 330-480 °C, 25-600 torr, respectively. The rate equations and appropriate Arrhenius rate constants for these three reactions are reported. The oxidation kinetics of C2F4 are also being investigated and preliminary results in the temperature and pressure ranges 175-300 °C and 25-200 torr are discussed. Finally the application of gas-solid chromatography techniques to the separation of low molecular weight fluorocarbons and the analysis of the C2F4 oxidation products are discussed.

#6006029
Mickle, E. A.; Clausion, W. B.; Szego, G. C.
FREE RADICALS AS HIGH ENERGY PROPELLANTS
General Electric Company, Flight Propulsion Laboratory, Cincinnati, Ohio
1956
106p
This compilation gives data concerning the use of free radicals as high energy propellants as well as discussions on rocket propulsion metastable states, and assumptions and methods of calculation. Tabulations and figures are included for the H, H2, F, F2, H2; N, N2, He; N, N2, H2; CH4, CH3, H; and F2, N2, H2 reactive systems. Included in the thermodynamic properties given for each system are: speed of sound in a gaseous mixture in which chemical equilibrium is maintained at all times, speed of sound in a gaseous mixture in which the chemical composition remains fixed, specific heat capacity at constant pressure for a gaseous mixture in chemical equilibrium, mole ratio of free radical to stable "parent" species in reactant mixture, mole ratio of diluent to stable parent species in reactant mixture, molal enthalpy, equilibrium constants, mass of a particle, molecular weight, specific or total entropy, specific volume, and alpha and beta emissions. Nozzle performance parameters and equilibrium composition in mole fractions are also listed. These data are listed for a broad range of temperatures and pressures for each system.

#6811003
Moiselwitzsch, B. L.; Smith, S. J.
ELECTRON IMPACT EXCITATION OF ATOMS
August 1968
120p
Available from Superintendent of Documents, GPO

The experimental and theoretical literature about the electron impact excitation of atoms is reviewed. Theoretical methods ranging from the Bennich-Born approximations to the close coupling approximations are discussed and intercompared. Where possible, on theoretical grounds or through intercomparison, the reliability of the various methods is discussed. A general critique of the optical method of measuring excitation functions is given, with the objective of promoting higher quality future experimental work. A critical study of existing experimental work leads to the conclusion that most work has failed to account for all important physical and instrumental effects, and it may be presumed that the data in the literature are subject to many unrecognized systematic errors. The literature on the alkali, heavy rare gases, mercury, cadmium, zinc is surveyed, but the quality of the literature does not support critical review beyond some general comments about the physics of these atoms.

#6609009
National Academy of Sciences, National Research Council. Committee for the Survey of Chemistry
CHEMICAL DYNAMICS - A CURRENT REVIEW
Publication 1292-B, National Academy of Sciences, Washington, D. C.
1966
66p
This report is a review of the current theory and experimental findings in chemical dynamics. It summarizes heterogeneous catalysis and electrochemical dynamics in surface chemistry, and includes reactions in solution as well as reactions in biological systems.

#7001042
Niles, F. E.
FORMATION OF IONS IN IONIZED AIR
BRL-R-1458, Ballistic Research Laboratories, Aberdeen Proving Grounds, Maryland
November 1969
54p
Available from DDC

Schematic representations are given for both the formation of positive ions and the formation of negative ions in ionized air. The formation of positive ions can be divided into three regions. Region I contains the ionization of neutral constituents and two-body reactions in dry air. Region II contains reactions involving ion species which are formed by three-body reactions. Region III contains reactions involving water vapor and shows the formation of the hydrated hydronium ions. The formation of negative ions proceeds predominantly through 0; the clustering of neutral molecules to 0; opens additional channels for the formation of NO3, which is not attacked by atomic oxygen. Rate constants are given, including the experimental values on which they are based, for the reactions in the schematic representations.

#6907039
O'Neal, H. E.; Benson, S. W.
THE BIRADICAL MECHANISM IN SMALL RING COMPOUND REACTIONS

The kinetics of dissociation, structural isomerization, and geometric isomerization in three- and four-membered ring compounds are discussed in terms of the biradical mechanism. With the exceptions of the cyclobutene isomerizations and the cis-1,2-alkylvinylcyclopropane isomerizations, all small ring compound reactions to date are shown to be both quantitatively and qualitatively consistent with the biradical mechanism. Methods for making a priori transition-state estimates of the activation energies and Arrhenius A factors for these reactions are outlined. Estimates are shown to be reliable to about 0.3 log unit in A and about 1.5 kcal/mol in E, on the average. The thermochromism of fluorine substitution in cyclopropanes and in cyclobutanes along with the effect of such substitution on the mechanism and kinetics of fluorinated small ring compound reactions is also discussed.

128
The Arhenius A factors for gas-phase, unimolecular reactions which proceed through a cyclic transition state are examined from the point of view of transition-state theory. The A factors are shown to be compatible with a structure which is looser than the ring prototypes. On the basis of a few simple rules for assigning bonding, stretching, and torsion frequencies it is shown that quantitative estimates of these A factors can be made to within the average experimental uncertainty of ±0.3 unit in log A (units of sec⁻¹). The extreme range of A values extends over 6.0 log units. In a few isolated cases the discrepancy between estimated and observed A factors exceeds 1.0 log unit and the weight of evidence suggests that the experimental values may not be reliable. The reactions include dehydrohalogenations which go through four-center transition states, ester pyrolyses, "ene" reactions, which go through six-center transition states, and Cope rearrangements which do the same. The analysis confirms previous suggestions that the most important contributions to ΔSᵗ come from losses in hindered internal rotations (about -4.0 g·cm²/rotor per rotor) in forming the loose cyclic structure. However, other coupled changes in structure can also be important.

Pitha, J.; Jones, R. N.
OPTIMIZATION METHODS FOR FITTING CURVES TO INFRARED BANDS
NRC Bulletin No. 12, National Research Council of Canada, Ottawa, Canada 1968 135p

Three computer programs for fitting infrared band contours are described and listed. These programs utilize Cauchy-Cratazer, Cauchy-Gauss, and Cauchy-Gauss product, and Cauchy-Gauss sum functions, and will optimize the parameters for fitting single bands or multiple overlapping band systems using a non-linear least squares algorithm. Supplementary programs are included for the evaluation of the half-widths and shape characteristics of the synthesized component bands, also for the computation of the coordinates of the synthesized band envelope, and for an analysis of the misfit with the true spectrum. This latter program also serves as a general purpose program for subtracting spectral ordinates. The programs are written in FORTRAN IV. They are designed for card input and output in a standardized format that facilitates their interfacing with other programs developed in our laboratory for the numerical analysis of infrared and ultraviolet spectrophotometric absorption curves.

Porter, G. (Editor)
PROGRESS IN REACTION KINETICS, VOLUMES 1 TO 4

This series reviews most of the principal branches of chemical kinetics and presents an up-to-date compilation of rate constants of current interest and importance. The chemical reactions are tabulated, rate constants given, the methods of measurement discussed, and the reliability of these values is assessed. The emphasis throughout is on the quantitative aspects of the subject and most of the articles contain original work not published elsewhere. The later volumes include cumulative indexes and extensions and amendments of the tables in previous volumes. The topics reviewed in the first volume range from fundamental reactions to combustion through diffusion kinetics to physical and chemical rate processes in solution and finally to biochemical kinetics. The articles in the second volume cover some of the major themes of chemical kinetics -- the reactions of halogen atoms and methylenes, mercury photosensitized reactions, anionic polymerization, cis-trans isomerization and protolytic reactions. Quantitative rate studies in biochemical systems are well illustrated by Gibson's account of haem reactions and the reaction index tabulates 1000 reactions for which rate data are given in the first two volumes of the series. The third volume begins with a theoretical review of homololytic reactions which includes an original discussion of some of the more obscure points of rate theory. There are articles on hydrogen atom reactions and hot atom chemistry and on the experimental technique of pulse radiolysis. Kinetic data obtained by this method are included. Gas-phase reactions are represented by a review of the fundamental physical process of vibrational relaxation and inhibition of radical chain reactions. A discussion of linear energy transfer in radiation processes and a review of recent work on the spectra and kinetics of aromatic radicals are given. There are two reviews of the kinetics of polymerization processes in solution, the first deals with polymerization by lithium alkyls and the second with the kinetics of radical polymerization processes. Volume four contains review articles on the reactions of organic and atmospheric radicals, dealing with halogenomethyl and alkoxyl radicals. Information is given on electronically excited states, including descriptions of recent progress made in this field. Proton transfer and kinetics of electrode processes are also discussed.

Porter, G. (Editor)
PROGRESS IN REACTION KINETICS, VOLUMES 5 TO 8

The later volumes include cumulative indexes and extensions and amendments of the tables in previous volumes. The topics reviewed in the fifth volume range from fundamental reactions to combustion through diffusion kinetics to physical and chemical rate processes in solution and finally to biochemical kinetics. The articles in the sixth volume cover some of the major themes of chemical kinetics -- the reactions of halogen atoms and methylenes, mercury photosensitized reactions, anionic polymerization, cis-trans isomerization and protolytic reactions. Quantitative rate studies in biochemical systems are well illustrated by Gibson's account of haem reactions and the reaction index tabulates 1000 reactions for which rate data are given in the first two volumes of the series. The third volume begins with a theoretical review of homololytic reactions which includes an original discussion of some of the more obscure points of rate theory. There are articles on hydrogen atom reactions and hot atom chemistry and on the experimental technique of pulse radiolysis. Kinetic data obtained by this method are included. Gas-phase reactions are represented by a review of the fundamental physical process of vibrational relaxation and inhibition of radical chain reactions. A discussion of linear energy transfer in radiation processes and a review of recent work on the spectra and kinetics of aromatic radicals are given. There are two reviews of the kinetics of polymerization processes in solution, the first deals with polymerization by lithium alkyls and the second with the kinetics of radical polymerization processes. Volume four contains review articles on the reactions of organic and atmospheric radicals, dealing with halogenomethyl and alkoxyl radicals. Information is given on electronically excited states, including descriptions of recent progress made in this field. Proton transfer and kinetics of electrode processes are also discussed.
This bibliography extends the JILA Information Center coverage of low energy atomic collision phenomena to include ion-molecule reaction rate data. It has been compiled by a search of earlier published literature for those papers that contain original experimental data on ion-molecule reaction cross sections or rate constants. The bibliography is divided into three parts. The first is a listing by reaction type under the Allen's headings. The second section is a list of references by citation number. The third section is an author index (all authors, not only the first).

**#6910046**

Sinnott, G. A.

**BIBLIOGRAPHY OF ION-MOLECULE REACTION RATE DATA**

JILA Report No. 9, Joint Institute for Laboratory Astrophysics, Boulder, Colorado

August 15, 1969

88p

This bibliography extends the JILA Information Center coverage of low energy atomic collision phenomena to include ion-molecule reaction rate data. It has been compiled by a search of earlier published literature for those papers that contain original experimental data on ion-molecule reaction cross sections or rate constants. The bibliography is divided into three parts. The first is a listing by reaction type under the Allen's headings. The second section is a list of references by citation number. The third section is an author index (all authors, not only the first).

**#6510037**

Somayajulu, G. R.; Zwolinski, B. J.

**A GENERALIZED BOND ENERGY SCHEME INCLUDING BARRIERS TO FREE ROTATION AND CORRECTIONS FOR ROTATIONAL ISOMERISM**

Chemical Thermodynamic Properties Center, Texas A&M University, College Station, Texas

1965

21p

A generalized procedure for predicting the energies of the paraffinic hydrocarbons is deduced on the assumption of (i) interactions between pairs of bonds attached to a carbon atom, (ii) interactions between trios of bonds attached to a carbon atom and (iii) interactions between pairs of bonds separated by a C-C bond. The derived equation justifies Tavestri's empirical approach and is shown to be not very different from Allen's equation without the steric terms. Under defined conditions this equation reduces exactly to Allen's equation. Barriers to free rotation were considered in deriving the new equation. This new equation is further modified by including corrections for rotational isomerism in place of trigonal interactions. The new relation is better founded on principle, correctly estimates enthalpies of atomization for normal and isomeric alkanes including those molecules without rotational isomers, and furthermore provides more refined steric terms due to higher order interactions.

**#6603012, 6603013, 6603014**

Somayajulu, G. R.; Zwolinski, B. J.

**RELATIONSHIP BETWEEN BOND ENERGY AND BOND DISSOCIATION ENERGY, II**

ENTHALPIES OF FORMATION OF THE SUBSTITUTED METHANES, SUBSTITUTED METHYL RADICALS AND SUBSTITUTED ETHANES,

III. MAGNITUDES OF THE BOND AND PAIR BOND CONTRIBUTIONS TO THE ENTHALPIES OF THE MOLECULES, IV. ENTHALPIES OF FORMATION OF SUBSTITUTED ETHYLENES

Chemical Thermodynamic Properties Center, Texas A&M University, College Station, Texas

August 1965; August 1965; September 1965

12p, 15p, 8p

In Part II, the enthalpies of formation of the substituted methanes have been calculated on the basis of Zahn's model. A method has been developed for the calculation of the dissociation energy of the R'-R" bond in a substituted ethane. Based on the dissociation energy of the R'-R" bond the enthalpies of formation of the substituted methyl radicals and the substituted ethanes have been calculated. The enthalpies of formation of the substituted methyl radicals can also be calculated on the basis of Zahn's model. In Part III methods have been developed for the calculation of the magnitudes of the bond and pair bond contributions based on relationships between bond energy and bond dissociation energy. The pair bond contributions for the pair of bonds C = in CX2, CX3, and CX4 have been determined. Also determined were the enthalpies of formation of CX2 and CS radicals, X being H, F, Cl, Br, I or CI, in Part IV enthalpies of formation of the CX3 type radicals obtained in Part III have been used to obtain the enthalpies of formation of the CX4 type radicals. Based on a relationship between bond energy and bond dissociation energy, the dissociation energies of the R'-R" bonds in substituted ethylenes have been calculated. Using the bond dissociation energies and the enthalpies of formation of the substituted methylene radicals, enthalpies of formation of a few substituted ethylenes have been calculated. The calculated values have been found to be in good agreement with the observed values.

**#6500415**

Stauffer, C. H. (Project Director)

**ALPHABETICAL INDEX TO TABLES OF CHEMICAL KINETICS HOMOGENEOUS REACTIONS**

NBS Circular S10, Supplement 2, National Bureau of Standards, Washington, D. C.

August 5, 1960

37p

Available from Superintendent of Documents, GPO

This Supplement to the Tables of Chemical Kinetics: Homogeneous Reactions, provides a subject index, an alphabetical index according to class of reaction, and an alphabetical index according to class of compound. As in Circular S10, each table is designated by a six-digit number, the first two of which refer to the type of reaction, the third to the name of the process, the fourth to gas, liquid, or solid. The second three-digit group of the table number refers to the types of substances involved.

**#6500414, 6500416**

Stauffer, C. H. (Project Director)

**TABLES OF CHEMICAL KINETICS - HOMOGENEOUS REACTIONS**

(SUPPLEMENTARY TABLES) VOLUME 1 AND VOLUME 2

NBS Monograph 34, Volumes 1 and 2 to accompany NBS Circular S10, Supplements 1 and 2, National Bureau of Standards, Washington, D. C.

Sept. 15, 1961, Volume 1; July 1, 1964, Volume 2

450p, 300p

Available from Superintendent of Documents, GPO

These tables form Supplement 3 to NBS Circular S10, Tables of Chemical Kinetics: Homogeneous Reactions, covering the period 1955-1959, for chemical reactions of the exchange-substitution and elimination type. The tables topics as a critically evaluated compilation of available factual numerical data on rates and rate constants obtained experimentally. The information compiled for each reaction is: amount of each reactant used and name of solvent; amount and name of catalyst used; if any, the rate constant, in terms of natural logarithms, temperature at which the measurement was taken, and a literature reference. For some reactions, the activation energy and Arrhenius constant are also given.

**#6906006**

Stevens, B.

**COLLISIONAL ACTIVATION IN GASES**


236p

The International Encyclopedia of Physical Chemistry and Chemical Physics is a comprehensive and current account of all aspects of the domain of science between chemistry and physics, written for the graduate and research worker. The subject matter is grouped in twenty topics. This third volume on the topic gas kinetics deals with energy...
transfer systems in gas reactions, relaxation techniques, rate measurements of competing processes, low and high energy studies, and higher vibrational level energy transfer. Tables give vibrational relaxation times and collision numbers for diatomic molecules; relative collisional deactivation probabilities for diatomic molecules M in presence of additive X; vibrational relaxation times, reduced relaxation times, and collision numbers for polyatomic molecules; and relative probabilities for collisional deactivation of polyatomic molecules by additives X.

#6500411, 6500413 *5.273
Thom, N. (Editor)
TABLES OF CHEMICAL KINETICS HOMOGENEOUS REACTIONS
NBS Circular 510 and Supplement 1, National Bureau of Standards, Washington, D. C., September 25, 1951; November 14, 1956
750p, 450p
Available from Superintendent of Documents, GPO

Circular 510 and Supplement contain a critically evaluated compilation of the available numerical data on rates and rate constants of homogeneous chemical reactions. The emphasis is placed on experimentally ascertained facts, and data depending on interpretations are not generally included. These tables represent a critically evaluated compilation of all the available factual numerical data on rates and rate constants of homogeneous chemical reactions. Stress is laid on experimentally ascertained facts. Data largely contingent on interpretations have generally not been included. In order to put the tabulations in uniform units, original data have been systematically recalculated wherever necessary. Reactions are arranged and numbered in the order of increasing complexity of the key reactants, thus in the case of organic compounds, in the order of increasing numbers of carbon atoms with brached isomers placed after the normal-chain isomers. Literature references and comments are appended to each table. Supplement 1 includes new tables, additions to the published tables, and revised tables canceling and replacing parts of the present tables. The tables are issued in the form of punched loose sheets for use in suitable loose-leaf binder.

#6809017 *5.271
Trotman-Dickenson, A. F.; Mline, G. S.
TABLES OF BIMOLECULAR GAS REACTIONS
120p
Available from Superintendent of Documents, GPO

This survey covers the kinetics of bimolecular and termolecular gas reactions that do not involve atoms or molecules in electronically excited states. Bimolecular reactions are here defined as reactions in which two molecules are involved as reactants, that yield two or more molecules as products. Those reactions in which two molecules combine to form one molecule are most usefully considered as the reverse of unimolecular reactions which will be dealt with in another survey. Reactions of oxygen and nitrogen atoms have been omitted as they will also form the subject of another survey. The literature surveyed covers the period from 1954 to December 31, 1965. The survey of earlier work has been based on one of the writers' books on "Gas Kinetics" which covered the literature to 1954. Use of the book for over ten years has revealed few omissions and these have been included in these tables. Data for the period January to August 1966 have been included where possible.

#6611001 *5.183-5.183
Veldre, V. I.
IONIZATION OF ATOMS BY ELECTRON BOMBARDMENT
Electron-Atom Collisions RICA 1965
100p
This is a review of the ionization of atoms by electron impact and of the theory of ionization. The article contains a bibliography of 103 references as well as 54 graphs and 5 tables of data. The article reviews the classical and quantum theory of ionization, the asymptotic behavior of the wave function, Born's Approximation, exchange calculations, calculation of distortion, momentum approximation, ionization from excited states, ionization of the helium atom and of complex atoms and ions, threshold behavior, and empirical formulas.

#6904006 *5.183
Venugopalan, N.; Jones, R. A.
CHEMISTRY OF DISSOCIATED WATER VAPOR AND RELATED SYSTEMS
New York: Interscience Publishers 1968
481p
This text reviews some of the latest studies and advances in knowledge of the oxygen-hydrogen system and explains experimental methods of dissociation, trapping, and analysis of dissociated and trapped products from water vapor and related systems. The book includes studies and reaction mechanisms for discharged water and hydrogen peroxide vapors, reaction of discharged hydrogen with oxygen and ozone, and for the products condensed at low temperatures from these systems. Thermal, photolytic, and radiolytic dissociations and mass spectrometric investigations of water and hydrogen peroxide vapors are summarized. The text describes reactions of hydrogen and oxygen pertinent to the chemistry of dissociated water vapor. It includes a discussion on superoxides of hydrogen and present knowledge of the methods of production of hydrogen and oxygen atoms and of hydroxyl and hydroperoxy radicals, their properties, and kinetic information on their reactions. Attention is focused mainly on gas-phase investigations.

#6905001, 6904014 *6.183-5.183
Warner, P. O.; Barry, H. F. (Compilers)
New York: Climax Molybdenum Company of Michigan, Inc. 1968, 1969
347p, 189p
The Molybdenum Catalyst Bibliography and Supplement 1 compile available references to over twenty different types of reactions involving various molybdenum catalysts. The broad types of reactions included in this compilation are hydroforming and reforming; cracking and hydrocracking; hydrotreating, hydrofining and desulfurization; hydrogenation; dehydrogenation; oxidation; reduction; amination and nitro reactions; hydrogenation, dehydrogenation, hydroxylation, hydroxylation and hydrolysis; carboxylation-carbonylation; Fischer-Tropsch synthesis; polymerization; isomerization; isotope exchange; condensation; chlorination; alkylation - dealkylation; catalyst preparation, activation, regeneration and physical properties; sulfurization-sulfonation; and others. Within each section the material is listed according to individual process and type of catalyst used and gives the appropriate reference for such information.
Colloid and Surface Properties
for single-element catalysts and alloys are included and interpreted, as are data illustrating differences due to the physical form of the catalyst (film, foil, wire, powder, and some supported systems). Problems are discussed concerning reproducibility of experimental results over repeated runs, and as a function of catalyst pretreatment. The data are analyzed in 29 and 29 tables, some of which are very extensive. The bibliography includes 141 references.

This book contains abstracts to 13,763 references which give data and information on solid adsorbents published primarily between 1943 and 1953. Some references to publications before 1943 have been added from the fields of heterogeneous catalysis and surface reactions on textile materials. The coverage is restricted to heterogeneous phenomena at solid-liquid and solid-gas interfaces. The patent literature is not covered. The book is divided into chapters on the general topics of adsorption of gases and vapors on solid adsorbents, adsorption from solutions on solid adsorbents, thermal effects in adsorption processes, theories of adsorption, refining of sugar and other applications of adsorbents, general information on adsorbents and special methods of investigation, and preparation of adsorbents. A subject index, author index, and list of commercial solid adsorbents are included.

This is the first volume in a series of monographs on physical chemistry designed as a reference source for experimental techniques, data, and ways to interpret results obtained in the study of monolayer behavior. The text includes discussions on the properties of liquid surfaces, experimental methods, properties of monolayer films, mixed monolayers, reactions in monolayers, transfer of monolayers to solids, multilayers, and scientific and technological applications. References to original reports are provided.

This volume of the Landolt-Bornstein tables contains data on fusion equilibria and interfacial tensions. The volume includes data from literature published through September 1955. Where several values for one measurement were found, the average was presented. Data on fusion equilibria are presented for binary and ternary systems of inorganic compounds, silicate systems, organic systems, and organic-inorganic systems. Interfacial tension data are presented for pure liquids towards their vapors or air, solutions towards air, two non-miscible systems, and interfacial films on water. Data are also presented on absorption from the gas phase and the liquid phase, and on paper chromatography.

This article is a tabulation of the cross-sectional area of gas molecules absorbed on solid surfaces. There are basically three ways to calculate these areas - molecular models, van der Waals constants derived from critical data, and liquid density, based on assumption of spherical shape and hexagonal close packing. In general, each method yields a different result for each molecule. The data compiled for each molecule includes a recommended value for the area, calculated by each method, followed by individual results calculated for different absorbents, together with the appropriate literature reference.
A bibliography of Liesegang rings (periodic precipitation figures produced by diffusion) and other periodic structures within the period 1855-1965 is listed alphabetically according to author. An author index and an index to applications of Liesegang rings to extra-chemical fields are also given. This second edition has added new references, which concern only true Liesegang rings, to the original bibliography of 681 references, many of which dealt with other kinds of periodic structures. The subject and author indices have been appropriately amended. Errors found in the first edition have been corrected. All journal article titles and nearly all book titles have been translated into English. Abbreviations of journal titles are those given in the issue of Chemical Abstracts in which the given reference first appeared.

The Molybdenum Catalyst Bibliography and Supplement compile available references to over twenty different types of reactions involving various molybdenum catalysts. The broad types of reactions included in this compilation are hydroforming and reforming; cracking and hydrocracking; hydrogenation, hydrofining and desulfurization; dehydrogenation; oxidation; reduction; amination and NH₃ reactions; hydration, dehydroxylation, hydrolysis; carbonylation-decarbonylation; Fischer-Tropsch synthesis; polymerization; isomerization; isotope exchange; condensation; chlorination; alkylation-dealkylation; catalyst preparation, activation, regeneration and physical properties; sulfurization-sulfonation; and others. Within each section the material is listed according to individual processes and type of catalyst used and gives the appropriate reference for such information.
Mechanical Properties
This is an authorized reprint from copyrighted publications of the American Society for Testing and Materials and the Specifications for Thermostat Metal Sheet and Strip (2-406-62); Methods of Test for: Flexibility of Thermostat Metals (2-106-56), Modulus of Elasticity of Thermocouple Metals (Cantilever Beam Method) (2-225-56), Maximum Loading Stress at Temperature of Thermostat Metals (Cantilever Beam Method) (2-305-56), Resistivity of Metallically Conducting Resistance and Contact Materials (2-406-54), Change of Resistance with Temperature of Metallic Materials for Electrical Heating (2-70-56), Mechanical Torque Rate of Spiral Coils of Thermostat Metal (2-362-62), Thermal Emissivity Rate of Spiral and Helical Coils of Thermostat Metal (2-389-62), Mean Specific Heat of Thermal Insulation (C 351-61), and Diamond Pyramid Hardness of Metallic Materials (E 92-57).

#6500221 #7.113
Beatty, G. H.; Ogden, H. R.
STATISTICAL ANALYSIS OF TENSILE PROPERTIES OF HEAT-TREATED Ti-4Al-3Mo-1V AND Ti-2.5Al-1V SHEET
DMIC Report 46J, Defense Metals Information Center, Battelle Memorial Institute, Columbus, Ohio June 6, 1961 42p
Available from CPSTI, P8 171 425

In connection with the Titanium Alloy Sheet-Rolling Program, DMIC has collected and processed tensile-property data submitted by producers of titanium sheet alloys. These data result from experimental production aimed at speedifying the development of high-strength, heat-treated sheet alloys for air frames and missiles. Ultimate strength, yield strength, and total elongation in 2" of Ti-4Al-3Mo-IV and Ti-2.5Al-1V are examined in this report. The sheet is produced by the Union Carbide Company (USC), Titanium Metals Corporation of America (TMCA), and Mallory- Sharon Metal Corporation (NS).

#7000103 #7.123-8.123
Campbell, J. E.
PLANE-STRAIN FRACTURE TOUGHNESS DATA FOR SELECTED METALS AND ALLOYS
DMIC Report S-28, Defense Metals Information Center, Battelle Memorial Institute, Columbus, Ohio June 1968 26p
Available from DDC

Plane-strain fracture-toughness (Kc) parameters may be used to estimate critical flaw sizes in structural metals subjected to known stresses at specified temperatures. Previous toughness parameters for evaluating high-strength alloys provided only empirical data that could not be used directly in design. This report contains the first compilation of available Kc data and is the result of considerable interest during the past several years in developing test methods for obtaining these data. The report is divided into sections on aluminum alloys, high-strength alloy steels, intermediate- and low-strength steels, precipitation-hardening stainless steels, titanium alloys, nickel-base alloy 718, and beryllium. Data on the aluminum alloys are limited to the -3000 and -7000-series alloys. The high-strength alloy steels include the maraging steels, 9Ni-4Cr steels, and martensitic steels such as AISI 430, 440, 403M, and H-11. The intermediate-strength steels include those that have been considered for submarine hulls, atomic-reactor vessels, and steam-turbine rotors. Data for the stainless steels are limited to the precipitation-hardening grades.

#6907030 #7.122
DATA STORAGE CONTENT OF THE MECHANICAL DATA FILE
Inventory Report 652, Mechanical Properties Data Center, Traverse City, Michigan January 1969 2p
Available from Mechanical Properties Data Center

This inventory presents an updated accounting of mechanical properties data and related information available from the Center. The data storage file consists of more than 750,000 mechanical properties test records, including associated material and test variables of more than 4000 structural metals and alloys. The chart is constructed to determine the quantity of retrievable test results by test type. Types of supplementary descriptive information and results are also tabulated.

#6500220 #7.123
Everhart, J. L.; et al.
MECHANICAL PROPERTIES OF METALS AND ALLOYS
NBS Circular C447, National Bureau of Standards, Washington, D. C. December 1, 1943 40p
Available from Superintendent of Documents, GPO

This circular is a summary of the results of a comprehensive survey of the technical literature on the strength and related properties, thermal expansion, and thermal and electrical conductivities of ferrous and nonferrous metals and alloys at temperatures over a wide range. In general, the data are presented in tabular form, although graphical representation is often useful to indicate the effects of changing composition or conditions on the properties. Data on aluminum, copper, silver, iron and steel, lead, nickel, tin, zinc, a number of miscellaneous metals and their alloys are included. The circular is not limited to conventional engineering materials but contains data on the properties of many materials not usually classed as such. Literature references to the sources of the data are included.

#6907029 #7.132
FATIGUE OF MECHANICAL AND WELD JOINTS OF TITANIUM 8-1-1
Search No. 2169, Mechanical Properties Data Center, Traverse City, Michigan (No date) 9p
Available from Mechanical Properties Data Center

This folder contains graphs for the fatigue of Ti-8Al-1Mo-V duplex annealed bimetal lap joint specimens for fatigue of Ti-8Al-1Mo-V annealed and welded sheet specimens for fatigue of Ti-8Al-1Mo-V duplex annealed and Ti weld 0.20" sheet specimens, and for the fatigue of Ti-8Al-1Mo-V duplex annealed and electron beam welded 0.20" sheet specimens plotting maximum stress versus cycles of life. Two pages of diagrams of fatigue specimens and a supplementary reference list are included.

#6700012 #4.137-7.133
Grigsby, D. L.
VANADIUM SILICIDE
Data Sheet BS-154 Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California April 1967 45p

These data sheets present a compilation of the electrical and electronic properties of vanadium-silicon compounds, especially VySi. These properties include transition temperature, specific heat and Debye temperature, critical field, electrical resistivity, penetration depth and absorption. Other properties included are: superconducting energy gap, magnetic hysteresis and susceptibility, critical current and current
density, Hall coefficient and thermoelectric power. Three additional sections of this report cover the crystallographic and mechanical properties of VSi as well as the nature of magnetic flux characteristics in the mixed state. Each of these properties is compiled over the widest possible range of parameters from references obtained in a thorough literature search.

#6907058

*7,033-8,033

Hamer, W. J.

RESUMÉ OF VALUES OF THE FARADAY


July-August 1968

A resume is given of the determinations of the value of the Faraday. Values obtained by silver deposition, iodide oxidation, oxalate oxidation, the omegatron, and silver dissolution are reviewed. All values are converted to the unified C12 international scale of atomic weights using the international atomic weights of 1967. Values of the Faraday are given in terms of both the NBS (legal) and absolute units of electrical measure. In the latter the new value for the acceleration due to gravity is used in computing the absolute value of electric current. On this basis and using the atomic weight of silver determined by Shields, Craig, and Dibeler, and converting to the C12 scale, the value of the Faraday is 96,486.9±1.6 absolute coulombs per gram-equivalent which differs by only 1 part per million from the value recommended by the National Academy of Sciences-National Research Council. If the atomic weight of silver recommended in 1967 by the International Atomic Weight Commission is used, the Faraday on the new gravity value is 96,486.541.6 absolute coulombs per gram-equivalent which differs by 5 parts per million from that recommended by the National Academy of Sciences-National Research Council. No change in the value of the Faraday adopted by the NAS-NRC Committee is recommended.

#6909032

*7,133-2,753

Hammond, J. P.

PHYSICAL, MECHANICAL, AND IRRADIATION PROPERTIES OF THORIUM AND THORIUM ALLOYS

ORNL-4219, Oak Ridge National Laboratory, Oak Ridge, Tennessee

April 1968

71p

Available from CPSTI

Physical and mechanical property data on thorium and thorium alloys were reviewed and information pertinent to nuclear reactor application is compiled.

#6906059

*7,143

Jurcovic, W. G.

STRUCTURAL PLASTICS APPLICATIONS HANDBOOK SUPPLEMENT I. TEXT METHODS


June 1969

45p

The procedures used for characterizing and evaluating structural plastics are described. Included are methods for determining physical, chemical, thermal, electrical, and mechanical properties.

#6500538

*0.1151-4.151-7.151-8.151

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME IV: TECHNOLOGY, PART 1: MATERIAL VALUES AND MECHANICAL PROPERTIES OF NON-METALS

Schmidt, E. (Editor)

Berlin, New York: Springer-Verlag

1955

88p

In German

This section of the Landolt-Bornstein tables contains material values and mechanical behavior of non-metallic substances, and their technical applications. Data are compiled from literature published through autumn 1964. The following general information is included: methods of systems, units and conversion tables, atomic weights, reduction to standard states, densities of water and mercury. Physical and mechanical properties of non-metallic solids are given for natural and synthetic building materials, organic natural materials (wood, paper, cellulose, wood pulp), fibrous materials, ceramics, glasses, synthetic materials, and natural synthetic materials. Also presented are data on friction and rolling resistance of non-metallic and metallic substances, viscosity, flow through tubes, continuous flow of bodies during two dimensional flow and rough flow, flotation, speed of sound, sound radiation and absorption, the human ear, sound spectra, vibrational numbers, spatial and building acoustics, sound recording and biological effects of ultrasonics.

#6500559

*0.1151-3.151-4.151-7.151-8.151

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME IV: TECHNOLOGY, PART 2a: MATERIAL VALUES AND PROPERTIES OF INDUSTRIAL METALS, BASIC PRINCIPLES, TESTING PROCESSES, INDUSTRIAL STEELS

Borschers, H.; Schmidt, E. (Editors)

Berlin, New York: Springer-Verlag

1963

88p

In German

This section of the Landolt-Bornstein tables includes data on metallographic principles and concepts, testing properties, and the properties of iron and its alloys. Data are given on diagrams of state, strength testing and destruction free materials. The physical properties of iron tabulated are lattice constants, density, thermal expansion, diffusion, atomic heats, specific heats, enthalpies, entropies and free enthalpies, phase diagrams and transition points, thermal conductivity, electrical properties, magnetic properties, optical constants and excitation cross-sections. Composition, treatment, strength values and special chemical properties of pure iron, unalloyed steels, highly alloyed steels, special steels, and cast iron are presented.

#6510029

*0.1151-4.151-7.151

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME IV: TECHNOLOGY, PART 2b: MATERIAL VALUES AND PROPERTIES OF INDUSTRIAL METALS, SINTERED INDUSTRIAL MATERIALS, HEAVY METALS (WITHOUT SPECIAL INDUSTRIAL MATERIALS)

Borchers, H.; Schmidt, E. (Editors)

Berlin, New York: Springer-Verlag

1964

1000p

In German

This section of the Landolt-Bornstein series deals with numerical data and behavior of both metallic and powder metallurgical industrial materials. The materials covered include the metals tungsten, tantalum, nio-
bium, vanadium, chromium, cobalt, nickel, manganese, gold, silver, copper, antimony, zinc, cadmium, lead, bismuth, tin, and their alloys. Data are provided for the physical and mechanical properties of these materials, their corrosion behavior and some of their applications to technology.

LANDOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY, GEOPHYSICS AND TECHNOLOGY, SIXTH EDITION, VOLUME IV: MATERIALS.

PART 2C: MATERIAL VALUES AND PROPERTIES OF INDUSTRIAL METALS, ALKALINE EARTHS, SPECIAL INDUSTRIAL MATERIALS, SEMICONDUCTORS, CORROSION

Borchers, H.; Schmidt, E. (Editors)
Berlin, New York: Springer-Verlag
1965
976p

In German

This section is the third part in a series of Landolt-Bornstein tables presenting the physical properties and behavior characteristics of most of the 75 elements in the periodic system exhibiting metallic characteristics as well as data concerning their alloys. Special emphasis is directed toward light metals, special work metals, binding materials, enamel properties, corrosion behavior, reactor metals, metals of the rare earths, semiconductors, and current developments in binary alloy systems. The volume is divided into sections entitled: titanium, beryllium, aluminum, magnesium, lithium, rubidium, cesium, liquid metals, general survey of reactor metals, urania, plutonium, zirconium, hafnium, thorium, rare earths, semiconductors, bearing metals, enamels, solder, welding and cutting metals, metal adhesives, and corrosive behavior of materials.

EXPONENTIAL MOLYBDENUM CHEMICALS

Larson, M. L.
Chemical Data Sheets EX-1 to EX-14, Chemical Data Bulletins Cub-1 to 13, Climax Molybdenum Company, New York, New York
March 1964-September 1966
200p

This is a collection of bulletins giving experimental data, physical properties, and references for various molybdenum compounds. The period covered is from March 1964 to September 1966.

MECHANICAL PROPERTIES OF STRUCTURAL MATERIALS AT LOW TEMPERATURES, A COMPILATION FROM THE LITERATURE

McClinstock, R. M.; Gibbons, H. P.
NAS Monograph 13, National Bureau of Standards, Washington, D. C.
June 1960
180p

Available from Superintendent of Documents, GPO

This compilation was taken from about 100 journal articles and technical reports on the mechanical properties of structural materials at low temperatures. It is published as an aid in the design of cryogenic equipment. The tensile strength, yield strength, tensile elongation and impact energy of about 200 materials, metallic and nonmetallic, are given graphically as functions of temperature between 4° and 300°K. The classes of materials considered are: aluminum, copper, nickel, titanium and their alloys; magnesium alloys; stainless steels, construction steels, super alloys, brazing and soldering alloys, miscellaneous alloys and pure metals, and nonmetallic materials.

The Materials Properties Data Book provides non-critical reference data of materials properties for use by persons involved with the analysis, design, fabrication or test responsibilities of the NERVA Project. The Data Book includes physical, mechanical and chemical properties of interest for materials of significance to the NERVA Program and, in addition, data on the effects of irradiation on materials. Information is also provided concerning material processing, with special emphasis on specifications. Material suppliers are given. The Data Book has been organized for the most effective use of the data by the design engineer. Volume I contains light metal alloys (aluminum, titanium, etc.) data. Volume I-A contains nickel-base alloys, refractory metals and alloys, and other nonferrous metals and alloys (cobalt, copper, etc.) data. Volume 2 contains alloys alloys data. Volume 3 contains nuclear materials (fuels, neutron controls, beryllium, etc.), graphite, refractory ceramics, adhesives and plastics data. The section for each individual material contains descriptive basic data sheets with pertinent data relative to properties, characteristics and other related information, together with curve sheets for graphic presentation of the parameters.

The Defense Metals Information Center has collected data generated in Phase I of the Titanium-Alloy Sheet-Rolling Program. The program is designed to develop the commercial production of four titanium alloys. The report presents the results of the analyses made to determine relationships between factors affecting the properties of the titanium alloy. The report is confined to the tensile properties of only two of the four alloys in the program. The report is divided into two parts. The first part examines the variance in tensile properties of the alloy. The second part examines some of the correlations between tensile properties and processing variables.

CURIUM DATA SHEETS

Rimshaw, S. J.; Ketchen, E. E.
ORNL-4186, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
27p

Available from CFSTI

Data on Cs-137 are tabulated. Properties of the fuel forms CsCl, Cs2SO4 and cesium borosilicate glass are given. Twenty-seven references are included.

CURIUM DATA SHEETS

Rimshaw, S. J.; Ketchen, E. E.
ORNL-4187, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
52p

Available from CFSTI

Data on the fuel forms of Cm-244 are presented. Properties of Cm-244 metal, Cm0.73, Cm0.92, Cm1.4,
Cm-242, 242CmO2 cermet are tabulated. Forty-nine references are given.

STRONTIUM-90 DATA SHEETS
ORNL-4185, Oak Ridge National Laboratory, Oak Ridge, Tennessee
December 1967
45p
Available from CFSTI

Data are given on composition, specific power, radiation, critical mass, container compatibility, thermophysical properties, mechanical properties, chemical properties, biological tolerances, and shielding for Strontium-90 metal, titanate (SrTiO3), oxide (SrO), fluoride (SrF2), and orthoflanate (Sr2F2O4). The data are indexed to 53 references which are also included.

*#6090007 *7.173-4.173 Kittelhouse, J. B.; Singletary, J. B.
SPACE MATERIALS HANDBOOK, THIRD EDITION
APML-TR-68-205, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio
July 1968
742p
Available from the DDC

This edition is the result of an extensive revision and development in line with the second edition of the Space Materials Handbook along with the incorporation of entirely new subject matter coverage and new materials data. All of the most significant material properties, phenomena, and principles covered in the original Handbook are presented and expanded in this revised and updated version. However, treatment of theoretical aspects has been considerably reduced, in order that more emphasis could be placed on the extensive new materials knowledge and data obtained from the design and successful launching of a wide variety of space systems. The Handbook is organized into four parts. Each part is presented in a space environment, effect of space environment on materials, materials in space and biological interaction with spacecraft materials. Information on mechanical, physical and chemical properties and characteristics is given for a wide variety of metallic and nonmetallic materials. The effects of natural and induced environments on materials are appraised. Materials categories include coverage of thermal control materials, optical materials, adhesives, organic structural materials, inorganic structural materials, electronic components and materials, materials for sealing applications and miscellaneous materials. A comprehensive multiple citation index is incorporated which gives ready access to information on specific subject areas with regard to their locations within the Handbook.

*#6050223, 6050224 *7.173-4.173 Schmidt, F. F.; Ogden, H. R.
ENGINEERING PROPERTIES OF TUNGSTEN AND TUNGSTEN ALLOYS, ENGINEERING PROPERTIES OF MOLYBDENUM AND MOLYBDENUM ALLOYS
DMC Reports 191 and 190, Defense Metals Information Center, Battelle Memorial Institute, Columbus, Ohio
September 1963; September 1965
135p
140p
Available from DDC

These two reports present the results of a state-of-the-art survey covering tungsten, molybdenum and 19 of their alloys. All data are given in tabular and graphical form covering some of the more important physical, mechanical and metallurgical properties for each material. For both reports, the physical properties included are melting point, density, thermal expansion, thermal conductivity and electrical resistivity. The mechanical properties included are tensile properties (ultimate tensile strength, yield tensile strength, elongation, reduction in area, modulus of elasticity) and their temperature dependence, notched tensile properties, creep and stress-rupture properties, and other selected mechanical properties. Metalurgical properties include: fabricability, transition temperature, weldability, stress-relief temperature and recrystallization temperature. The ten alloys considered in the tungsten properties report include "doped" tungsten and various alloys with niobium, thorium dioxide, molybdenum and rhenium. The nine alloys considered in the molybdenum properties report include alloys with niobium, zirconium, titanium, tungsten and carbon. References are given at the conclusion of each materials section.

*#6090025-6090028 *7.173 Schwartzberg, F. R.; et al.
CRYOGENICS MATERIALS DATA HANDBOOK
Technical Documentary Report No. ML-TDR-66-280 and Supplements 1, 2, 4, 5; The Martin Company, Denver, Colorado
400p, 150p, 150p, 150p
Available from CFSTI; AD 609 562, AD 611 165, AD 618 065, AD 633 388

This is a revision of the previously published "Cryogenic Materials Data Handbook" and its supplements with the CFSTI designation PB 171 809. It is organized into sections of aluminum, stainless steel, titanium, superalloys, alloy steels, miscellaneous metals, alloys, polymeric materials and fiber-reinforced plastics containing mechanical and physical property data on 62 metallic and nonmetallic materials. The properties measured include yield, tensile, notched tensile, weld tensile, impact, compressive, fatigue, shear and flexural strengths; elongation; reduction of area; fracture toughness; stress-strain diagram; modulus of elasticity; and rigidity; hardness; compressive, shear and flexural moduli; thermal expansion; Poisson's Ratio; thermal conductivity; and resistivity. The Handbook contains material, property and accumulative data and a reference list. Progress reports are issued semiannually to update and maintain the Handbook. The first Supplement to the Cryogenic Materials Data Handbook updates the publication to February 1965. The subsequent supplement updates it to July 1965, and the last to March 1966.

*#690061 *7.173-4.173 Seddon, B. J. (Compiler)
PHYSICAL PROPERTIES OF SOME PLUTONIUM CERAMIC COMPOUNDS: A DATA MANUAL
TRG-Report 1601, United Kingdom Atomic Energy Authority, Reactor Group, Risley, England
December 1967
54p
Available from CFSTI

This report is a compilation of the best available data on the physical properties of some of the plutonium compounds which are, or may be, of interest as fuel materials. These are the dioxide, monocarbide, mononitride, monosulfide and monophosphide of plutonium, uranium-plutonium dioxide, uranium-plutonium monocarbide, uranium-plutonium mononitride, and the plutonium silicides.

*#6907032 *7.171 SELECTED BIBLIOGRAPHY, MECHANICAL PROPERTIES OF Ti-6Al-4V REVISIRED,
Chapter 3707, Ti-6Al-4V Revised, Mechanical Properties Data Center, Traverse City, Michigan
1969
102p
Available from Mechanical Properties Data Center

This is a set of computer printout sheets giving the titles and sources of selected references to the literature for data and information on the mechanical properties of Ti-6Al-4V.
This report contains computer printout sheets of data on the tensile properties of Udiment 700 at room and elevated temperatures including the primary operation, secondary operation, heat treatment, specimen configuration, specimen thickness or diameter, notch configuration, orientation, test environment amount, ultimate tensile strength, yield stress, gage length, percent elongation, percent reduction in area, along with the reference source.
Thermodynamic and Transport Properties
This monograph deals with the semiconducting compounds formed from elements of groups II, IV, or V and sulfur, selenium, or tellurium. The monograph describes the crystal structure of these compounds and presents their physical properties on their polymorphic transitions, including transitions observed at high pressures. The monograph includes published P-T-X diagrams for the compounds and discusses a method for plotting the diagrams. The more important physicochemical properties of the compounds are also given: the forbidden band width, the carrier mobility, the effective masses of carriers, the electrical conductivity, and the thermoelectric power. The monograph consists of three chapters. The first deals with chalcogenides of elements in group II: zinc, cadmium, and mercury. The second discusses semiconducting compounds of elements in group IV: germanium, tin, and lead. And the third describes investigations of compounds of elements of group V: arsenic, antimony, and bismuth. The monograph does not cover chalcogenides in group III. Also excluded are data on semiconductor glasses. The monograph includes the published data up to the second half of 1966 on the crystal structure, physicochemical, and electrical properties of the phase diagrams of important semiconducting compounds which are used in various branches of engineering.

This is a translated version from the Russian.
(Cantilever Beam Method) \( (\text{B} 305-56) \), Resistivity of Metallically Conducting Resistance and Contact Materials (\( \text{B} 63-49 \)), Change of Resistance with Temperature of Metallic Materials for Electrical Heating (\( \text{B} 70-56 \)), Mechanical Torque Rate of Spiral Coils of Thermostat Metal (\( \text{B} 362-62 \)), Thermal Deflection Rate of Spiral and Helical Coils of Thermostat Metal (\( \text{B} 389-62 \)), Mean Specific Heat of Thermal Insulation (C 351-61), and Diamond Pyramidal Hardness of Metallic Materials (C 92-57).

# American Society for Testing Materials

**ASTM VISCOSITY INDEX CALCULATED FROM KINEMATIC VISCOSITY**


852p

The tables in this publication permit direct reading of the Viscosity Index of a petroleum or lubricant if its kinematic viscosities at 100 F and 210 F are known. Use of these tables eliminates the calculation required for determining Viscosity Index.

# American Society for Testing Materials Committee D-2 on Petroleum Products and Lubricants

**PHYSICAL CONSTANTS OF HYDROCARBONS C-1 TO C-10**


60p

The tabulation furnishes constants in both the English engineering and metric systems of measurement. The major sources of data for this tabulation are the findings of Research Project 44, Data on Hydrocarbons and Related Compounds, and Research Project 45, Synthesis, Purification, and Properties of Hydrocarbons of Low Molecular Weight, of the American Petroleum Institute. These data are either critically selected, precisely calculated or deemed to be of good experimental quality. Data included from other sources are shown in italics. Data calculated for this compilation are shown in parentheses. The atomic weights used are based on oxygen = 16.000, carbon = 12.01, and hydrogen = 1.008. The recent changes in the atomic weight scale are so small that for engineering use the change in the atomic weight scale is not important.

# Aroeste, H.; Magee, J. L.

**THERMAL RADIATION PHENOMENA, VOLUME 3. TABLES OF RADIATIVE PROPERTIES OF AIR**


635p

Available from CFSTI, AD 564773

This volume presents tables of the equilibrium radiant properties of air and its constituents for a wide range of temperatures and densities. The work is divided into five parts: Part A is a summary section designed to give an overview of the field. Some of the data are presented in semi-schematic form and should not be used for quantitative work. Part B gives basic data and references used in the preparation of the subsequent parts. Finally, parts C, D, and E contain the absorption coefficients in the ranges 1000°C to 74,000°C (Part C), 1 eV to 20 eV (Part D), and above 20 eV (Part E).

# Bain, R. W.

**STEAM TABLES 1964 PHYSICAL PROPERTIES OF WATER AND STEAM 0-300°C 0-1000 BARS**

Edinburgh, Scotland: Her Majesty's Stationery Office, 1964

147p

These tables, prepared at the National Engineering Laboratory of the Department of Scientific and Industrial Research, provide physical properties of water and steam. The tables are divided into three parts: Table 1 contains values of the saturated liquid and vapor at round values of temperature, Table 2 contains the same quantities at round values of pressure. Table 3 contains values of the specific volume, specific enthalpy and specific entropy of both liquid and vapor at equally spaced intervals in the temperature range 0-300°C and at varyingly spaced intervals in the pressure range 0-1000 bars. The international units used are those accepted for the international skeleton tables.

# Ballard, R. E.; Nelson, E.

**ABSORPTION OF PLANTED SOLID DRUGS**


Factors that clearly affect absorption rates of implants are their surface area and solubility in body fluids. Factors proposed that do not affect absorption rate greatly (if at all), with the present information, are pellet density (when area corrections are made), crystal size used in implant preparation, phagocytosis, physiological need or sex of the animal, encapsulation, "ghost" formation, and the age of the animal (if it is not young). Factors such as site of implantation and body movement and diluents do have an effect, but little quantitative information is available concerning the magnitude of the effect.

# Ballard, S. S.; McCarthy, K. A.; Wolfe, W. L.

**OPTICAL MATERIALS FOR INFRARED INSTRUMENTATION**

IRIA Report 2339-118, Michigan University, Willow Run Laboratories, Ann Arbor, Michigan, January 1959

115p

Available from NDC, AD 217367

The properties of approximately fifty materials which are useful in optical instrumentation in the infrared portion of the electromagnetic spectrum are described and compared. Each material is discussed in detail in terms of its optical, chemical, thermal, and mechanical properties. Tables and figures are given so that the properties of the various materials can be readily compared.

# Ballard, S. S.; McCarthy, K. A.; Wolfe, W. L.

**OPTICAL MATERIALS FOR INFRARED INSTRUMENTATION, SUPPLEMENT**

Report 2339-11-8, Michigan University, Infrared Laboratory, Ann Arbor, Michigan, April 1961

10p

Available from CFSTI, AD 255699

In this supplement, additional data are compiled on the optical materials, which are useful for infrared instrumentation, treated in 2339-11-8, as well as data on three new materials, Irtran-1, Irtran-2, and diamond. In addition, several errors and ambiguities in the original report are corrected and clarified.
This paper is a review of seven hundred and ten articles on low-temperature phenomena published during the period from 1763 to 1948. A historical review is given: the main calorimeter types are described briefly; and the thermal data on all substances measured have been collected in tabular form. Author and formula indexes have been constructed.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

Data relating to the isentropic one-dimensional expansion of an ideal gas with a constant ratio of specific heats are presented in tabular form. In Part I, the tables are calculated for the value γ = 1.150.

This text, translated from Russian, is a review of the properties of silicon and its binary compounds. The text discusses the preparation of binary silicon compounds, properties and uses of H-Si and Halogen-Silicon systems, systems formed by silicon with elements of Groups IA, IB, II A, II B, III A, III B, IV A, VII A, VII B, transition metals of Groups IV, V, VI, VII, and VIII, and with metals of the iron subgroup, palladium subgroup, platinum subgroup, lanthanides and actinides. Ternary and quaternary systems are also included. There is a list of 716 references and tables of data accompany the systems discussed.

This bibliography contains 452 references of low temperature, specific heat data and information for elements, compounds or alloys, of all elements having atomic numbers 1 to 84, 90, 92, and 94, with the exception of the inert gases and technetium. The bibliography is accompanied by an index of elements and compounds.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.

In order to develop some preliminary information concerning the very basic problems of gaseous deposition at extremely low temperatures, samples of argon, nitrogen, oxygen, and hydrogen were deposited at 4.2 K, and variations in some of the parameters associated with these depositions were observed. In addition, samples of the same gases were passed through an excitation zone prior to deposition. The differences in the deposition behavior were recorded. The pressure downstream of the deposition region and the temperature within the deposited solids were found, in general, to increase with flow rate of the depositing substances, and to vary with time in a manner depending on the density, flow rate, and state of excitation of the incoming gases. Arrhenius curves were plotted using warming data observed in these experiments and solid-phase transition data determined elsewhere. Variations in the positions of these plots were sometimes the result of differences in the conditions of deposition, and at other times an effect of the chemical nature of the gas studied. The experimental behavior of hydrogen, relative to that of the other gases studied, was observed to be strongly dependent upon its much higher thermal conductivity.
Equilibrium thermodynamic data are presented for argon from a temperature of 100 to 15,000 K and from a log ρ of -7.0 to 3.0, where ρ is density in anagats. These data are tabulated at constant temperature with density incremented. The dependent parameters are log ρ, log P, log Hz/RT, gamma, A/AA, 2, Hz/RT, and γ*. Several sources were used in assembling the data.

Equilibrium thermodynamic data are presented for nitrogen from a temperature of 100 to 15,000 K and from a log ρ of -7.0 to 3.0, where ρ is density in anagats. These data are tabulated at constant temperature with density incremented. The dependent parameters are log ρ, log P, log Hz/RT, gamma, A/AA, 2, Hz/RT, and γ*. Several sources were used in assembling the data.
Equilibrium thermodynamic data are presented for nitrogen from an S/R of 11.6 to 135.0 and from a temperature of 100 to 15,000 K, where S/R is dimensionless entropy. These data are tabulated at constant entropy with temperature incremented. The dependent parameters are log $P$, log $z$, log $U/R$, gamma, gamma, $A/AA$, $Z$, $H/R$, and $Z^*$. Several sources were used in assembling the data.

Table 7.11.3

<table>
<thead>
<tr>
<th>Substance</th>
<th>Density</th>
<th>Heat Capacity</th>
<th>Entropy</th>
<th>Heat Capacity</th>
<th>Entropy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrogen</td>
<td>1.23</td>
<td>1.50</td>
<td>0.50</td>
<td>1.75</td>
<td>0.85</td>
</tr>
</tbody>
</table>

A systematic compilation of a bibliography on heat-capacity data of ammonia, carbon dioxide, and water is being made. The literature data examined thus far are listed. The list is probably 30 percent complete.

Other toughness parameters may be used to estimate critical flaw sizes in structural metals subjected to known stresses at specified temperatures. Previous toughness parameters are used for evaluating high-strength alloys. Data on the aluminum alloys are limited to the 2000- and 7000-series alloys. High-strength alloy steels include the maraging steels, 09M-400 steels, and alloy steels such as AISI 4340, 6625, 300N, and H-11. The intermediate-strength steels include those that have been considered for submarine hulls, atomic-reactor vessels, and steam-turbine rotors. Data for the stainless steels are limited to the precipitation-hardening grades.

The bibliography lists about 1500 references on the equation of state of materials at high pressure. These are listed chronologically; each citation is accompanied by descriptors for the materials under theories investigated. The descriptors have been indexed alphabetically according to the "keyword" system. Thus it is possible to locate references dealing with any substance or theory, or with various other keywords, by looking in the keyword index (Part 1), and then finding the reference itself in the bibliography (Part 2) by its serial number. The indexing is more complete for elements, inorganic compounds, minerals, and explosives than for organic compounds and composite materials. Most mixtures have been excluded. For gases, "high pressure" is taken to mean pressures above 50 atm, or virial coefficients higher than the second. For liquids and solids, all compressibility measurements are included when found. Topics such as shock-wave compression, melting curves, quantum theory of the electron gas, and Thomas-Fermi theory have also been included.
It is shown that the Principle of Corresponding States is applicable to thermal conductivity of metals. Two correlative relations are presented: one at low temperatures \(0 < T < 1.5T_m\) and another at moderate and high temperatures \(0.5T < T < 3.0\), where \(T_m\) is the temperature corresponding to the thermal conductivity maxima and \(\theta\) is the Debye temperature. A universal relation for the reduced thermal conductivity is obtained in the low temperature region based on the solid state physical theory of metals and the proposed Principle of Corresponding States.
This report presents a table of thermodynamic properties of gaseous hydrogen covering the temperature range from 100 to 3000°K and the pressure range from 1 to 50 atmospheres. The calculations have been done by means of a computer routine. The table has been made for temperature at 10° increments. Pressures are at 1 atmosphere and at 5 atmosphere increments from 5 through 50 atmospheres, and for 25%, 50%, 75%, and 100% of the gas initially in the para state. The first of these latter values corresponds to normal hydrogen, whereas the remaining values apply to vapor from liquid which has been stored for varying lengths of time. For each entry, the table gives the density in gm/liter, the entropy in joules/gm, the entropy in joules/(gm°K) and the fraction of the gas molecules which are dissociated.

Cohn, E. J.; Edsall, J. T.
PROTEINS, AMINO ACIDS AND PEPTIDES AS IONS AND DIPOLAR IONS
404p
In this book an attempt has been made to characterize amino acids, peptides, and proteins. The evidence concerning the size and shape of these molecules and the number and distribution of the ionic charges on them has been examined. The implications of the molecules' charged structure by their physical properties, and the physicochemical interaction with other molecules is considered. It is not a treatise on all aspects of the chemistry of the proteins, or of their physical chemistry, but is aimed at subjects which are of primary concern. Experiments relating to the physical properties of the proteins have been described in which it has been necessary to do so in order to make clear the operational basis on the principles which are involved.

Collins, F. C.
THEORETICAL AND EXPERIMENTAL STUDIES OF LIQUID VISCOITY
AADC Technical Report 57-413, Aeronautical Research Laboratory, Wright-Patterson Air Force Base, Ohio
July 1957
35p
Available from DDC, AD 139871
A review of the theoretical background is presented together with analysis of the outstanding difficulties of the various theories of liquid viscosity. The present status of the rigid sphere model of liquid viscosity and its relationship to sonic velocity is reviewed. The experimental data relating to the viscosity and sonic velocity in a number of typical liquids are presented and tentatively evaluated in terms of the rigid sphere model of liquid viscosity. A brief description is given of the apparatus for measuring sonic absorption in liquids.
The tabulation is based on papers and reports received by the NRC through January 1960.

**CONSOLIDATED INDEX OF SELECTED PROPERTY VALUES, PHYSICAL CHEMISTRY AND THERMODYNAMICS**


275 p. Available from Superintendent of Documents, GPO

The Consolidated Index is a key to the contents of publications that present critically evaluated numerical property values. This initial volume contains in codified form the contents of the following six compilations of physicochemical and thermodynamic data: Selected Values of Properties of Hydrocarbons and Related Compounds (American Petroleum Institute Research Project 43), Selected Values of Properties of Chemical Compounds (Manufacturing Chemists Association Research Project), Selected Values of Chemical Thermodynamic Properties of the Elements (D.R. Stull and G.C. Sinke), Contributions to the Data on Theoretical Metallurgy (Bureau of Mines Bulletins 385, 384, 395, 396, 407, 477, 524, 584), and Selected Values for the Thermodynamic Properties of Metals and Alloys (Minerals Research Laboratory, University of California). Each element, compound, or system of properties is accessed as a separate index entry. For a given substance, all properties that appear in one or more of the compilations are listed separately under each. A given property, if appearing in several, is repeated under each publication.

**ELECTROCHEMICAL DATA**

Conway, B.E.


374 p.

A series of more than 300 tables of electrochemical data selected from the best quantitative information available is presented. The range of subject matter includes universal constants, properties of the rare gases, biological importance, conductivities and potentials in fused salts, and parameters of electrode reaction kinetics, as well as the more usual compilations of conductances, standard electrode potentials and transport numbers. The author is meticulous in giving references to the original sources and his extensive bibliography may be almost as useful as the data themselves. Whenever possible he states the precision of the values given and in many cases indicates the experimental or theoretical method by which they were obtained. The hook is well organized, beginning with universal constants and general physical properties progressing through the thermodynamics of solutions to ionic transport. The author then deals with the electric double layer at interfaces. After a diversion into the field of fused and solid ionic conductors, he returns to the logical order with a chapter on reversible electrode reactions. The concluding section, of which Dr. J. O'M. Bockris is the author, is an attempt to collect data on electrode kinetics, giving tables of the Tafel parameters. Most of the material deals with hydrogen evolution but there are tables on the deposition of metals, redox processes and anode reactions such as the evolution of oxygen and the halogens. There is a 14-page index to provide ready reference to specific information.

**COOPER, J. (Editor) PROCEEDINGS OF WORKSHOP CONFERENCE ON THE LOWERING OF THE IONIZATION POTENTIAL AND RELATED PROBLEMS OF THE EQUILIBRIUM PLASMA**

JILA Report No. 79, Joint Institute for Laboratory Astrophysics, Boulder, Colorado July 1966

225 p.

This workshop was sponsored by the Joint Institute for Laboratory Astrophysics. Its concern was statistical mechanics of plasma both from experimental and theoretical points of view. There were many recent theoretical advances in the treatment of such problems as the plasma partition function and the lowering of the ionization potential, and the organizers' wish was to bring together active theorists and experimenters in this field to resolve some of the differences. Included in the Appendix is a Bibliography on Research on Lowering of the Ionization Potential.

**COOPER WIRE TABLES**


41 p. Available from Superintendent of Documents, GPO

This handbook is a revision of the Copper Wire Tables previously published as NBS Circular 31. It reflects changes in the nominal diameters of gages 45 and smaller and extends the tables to 56 gage. The handbook provides data for the weight and length per ohm, weight per unit length, length per unit weight of copper wire, resistivity, temperature coefficient, density, and conductivity of copper wire from 0000 to 56 gage as a function of temperature from 0 to 290°C.

**CORRUCCHINI, J.R.; GNAWEK, J.J. SPECIFIC HEATS AND ENTHALPIES OF TECHNICAL SOLIDS AT LOW TEMPERATURES, A COMPILATION FROM THE LITERATURE**


20 p. Available from Superintendent of Documents, GPO

A compilation of heat capacity data on cryogenic materials for designers of cryogenic equipment is given. Tables are given of the specific heat and the enthalpy of 98 metals, 3 alloys, 8 other inorganic substances and 8 organic substances in the temperature range 1° to 300°K. The data presented are based on selected sources listed under each table, including references which did not influence the tabulated data but are of current interest. The 170 references cover the period 1930 to 1960. The research was based on Physics Abstracts and Chemical Abstracts for those years not covered by bibliographies of Kelley (1950) and Schiffman (1952).

**CORRUCCHINI, J.R.; GNAWEK, J.J. THERMAL EXPANSION OF TECHNICAL SOLIDS AT LOW TEMPERATURES, A COMPILATION FROM THE LITERATURE**


22 p. Available from Superintendent of Documents, GPO

Tables are given of the linear contraction relative to 293°K (L293 - L) of thirty elements, forty-
five alloys, twenty-two other inorganic substances and twenty plastics and elastomers in the temperature range, 0 to 300°C.

Coughlin, J. P.
CONTRIBUTIONS TO THE DATA ON THEORETICAL METALLURGY XII. HEATS AND FREE ENERGIES OF FORMATION OF INORGANIC OXIDES
80p
Available from Superintendent of Documents, GPO

This bulletin compiles heat and free energy of formation data for inorganic oxides. A survey of worldwide literature to September 1953 permitted assembly of data for over 170 inorganic oxides. These data are presented by three separate methods: (1) tables giving heats and free energies of formation values at 298.16 K, at phase-change temperatures and at even 100° intervals from 400° to a maximum of 2000°K, (2) equations giving free energies of formation as functions of temperature, utilizing in their derivation the best available high temperature heat-content values, and (3) equations of more approximate nature giving free energies of formation as linear functions of temperature.

Cryogenic Data Center, National Bureau of Standards, Boulder, Colorado
1965
30p

These cryogenic data memoranda contain thermodynamic data and reference sources for properties of gas−solid vapor pressure, equilibrium deuterium heat of vaporization, normal deuterium saturation density of liquid, solid helium density, and composition of ethane, butane, pentane density and compressibility, and solid hydrogen density and compressibility.

Darwent, R. deB.
BOND DISSOCIATION ENERGIES IN SIMPLE MOLECULES
January 1970
$2p
Available from the Superintendent of Documents, GPO

Bond dissociation energy values (kcal/mol) and (kJ/mol) of simple compounds are tabulated from a literature review covering the years 1962−1966 inclusively. Some selected values which appeared in the years 1956−1962 are also included. Organic compounds are excluded except those containing one carbon atom. The groups −CO and −CN are not considered to be organic. The values are quoted usually at 0°K or 298°K and refer to the gaseous state. They represent the energy required to break a bond at the specified temperature with all substances in the zero vibrational state of the ground electronic state. The experimental method for the energy value listed is given and referenced in the table. A value recommended by the author is listed as the final value for each reaction.

Dean, J. V.
A TABULATION OF THE THERMODYNAMIC PROPERTIES OF NORMAL HYDROGEN FROM LOW TEMPERATURES TO 300°K AND FROM 1 TO 100 ATMOSPHERES
NBS Technical Note 120, National Bureau of Standards, Boulder, Colorado
November 1961
71p
Available from the Superintendent of Documents, GPO

Pressure, volume, temperature, internal energy, enthalpy, and entropy of normal hydrogen gas have been tabulated along isobars in 1°K temperature steps. The range covered is from the saturation temperature to 300°K and from a pressure of 1 to 100 atmospheres. The source of data is the Research Paper 1932 of the National Bureau of Standards Journal of Research. The method is described by which the data presented in Research Paper 1932 is reduced to properties directly useful for engineering calculations. A method is also described for estimating the effect of ortho−para compositions upon the tabulated properties. Tabular values are presented in the dimensional units of the metric system. The tabulations are also available in the dimensional units of the British system in Supplement A of the Technical Note 120.

Dean, J. V.
A TABULATION OF THE THERMODYNAMIC PROPERTIES OF NORMAL HYDROGEN FROM LOW TEMPERATURES TO 540°R AND FROM 10 TO 1500 PSIA, SUPPLEMENT A (BRITISH UNITS)
NBS Technical Note 120A, National Bureau of Standards, Boulder, Colorado
June 1962
75p
Available from the Superintendent of Documents, GPO

Pressure, volume, temperature, internal energy, enthalpy, and entropy of normal hydrogen gas have been tabulated along isobars in 2°K temperature steps. The range covered is from the saturation temperature to 540°R and from a pressure of 10 to 1500 psia. The source of data is Research Paper 1932 of the National Bureau of Standards Journal of Research. The method is described by which the data presented in Research Paper 1932 is reduced to properties directly useful for engineering calculations. A method is also described for estimating the effect of ortho−para compositions upon the tabulated properties. Tabular values are presented in the dimensional units of the British system. The tabulations are also available in the dimensional units of the metric system as Technical Note 120.

Dennison, D. H.; Gschneidner, K. A.
A TABULATION OF THE SPECIFIC HEAT CONTRIBUTION DUE TO THERMAL EXCITATION (SCHOTTKY EFFECT) OF THE 4f ELECTRONS OF SOME DI- AND TRI-VALENT LANTHANIDE ELEMENTS
IS-1156, Ames Laboratory, Ames, Iowa
May 1965
37p
Available from CFSTI

The 4f electronic specific heat contribution for divalent samarium and for the trivalent lanthanide elements Ce, Pr, Nd, Sm, Eu, Gd, Dy, Ho, Er, Tm and Yb has been calculated and is tabulated at 25°K intervals from 50° to 2000°K at 50°K intervals from 200° to 1000°K, and at 100° intervals from 1000° to 2000°K.

Diamond, J. J. (Editor)
BIBLIOGRAPHY ON THE HIGH TEMPERATURE CHEMISTRY AND PHYSICS OF MATERIALS, October, November, December 1968; January, February, March 1969; April, May, June 1969; July, August, September 1968; October, November, December 1969
NBS Special Publication 315, 315-1, 2, 3 & 4, National Bureau of Standards, Washington, D. C.
April, July, October 1960; June, 1976
72p, 31p, 82p, 90p, 35p
Available from Superintendent of Documents, GPO

These bibliographies consist of references to research involving temperatures above 1000°C, which were noted by the contributors during the above-mentioned three-month periods. Since these are intended primarily as current-awareness

159
Thermochromical research was carried out on ethylenimine, four aliphatic diamines, and two organic fluorne compounds. Results for ethylenimine include an experimental determination of its thermodynamic properties by means of low-temperature calorimetry, combustion calorimetry, and vapor pressure measurements. The measured third-law entropy is discussed in light of a recent Russian value calculated from spectroscopic information. Work on the thermochrometry of aliphatic diamines includes purification of samples of 1,2-diaminoethane, 1,2-diminoopropane, 1,2-diminoobutan e, and 1,2-dimino-2-nilethylpropane; heats of combustion and vapor pressures of 1,2-dimino-2-nilethylpropane; and some low-temperature thermal measurements on 1,2-diminoopropane. P-V-T relations determined for hexafluoroethane include vapor pressures to the critical state. Critical constants, $P_c$, $T_c$, and $V_c$, and compressibility of the gas up to $350°C$ and $400$ atmospheres. Preliminary values for the vapor heat capacity of hexafluorobenzene are presented.

Vapor-liquid equilibria data for the hydrogen-methane system are presented in the form of K-factor charts for thirteen isotherms from $90.3$ to $188.2°C$ and from $30$ to $270$ atmospheres pressure. Solid-vapor equilibria data are presented as vapor phase compositions versus temperature and pressure over the range of $57$ to $90°C$ and $5$ to $15$ atmospheres. A bibliography of 290 references on related properties for this system and for the pure components is also presented.

Literature data have been used to calculate K-factors for the hydrogen-carbon monoxide system over the range of $68.2$ to $122.2°C$ and for 10 to 225 atmospheres pressure. K-factors are presented graphically for each isotherm over this range. Published data on the solid-vapor region are presented separately as composition versus pressure at constant temperature. A bibliography of approximately 450 references is also presented on related properties for this system and for the pure components.
Published data have been used to calculate K factors for the hydrogen-nitrogen system over the liquid-vapor range of 68.2 to 122.2°C, and 10 to 275 atmospheres. K factors are presented graphically for eleven isothersms within this range. Published data on the solid-gas and solid-liquid regions are presented separately as composition versus pressure at constant temperature. A bibliography of 250 references pertaining to the hydrogen-nitrogen system is included.

#6500324 1964
Drayer, D. G.
THE VAPOR PRESSURES OF 30 INORGANIC LiquIDS BETWEEN ONE ATMOSPHERE AND THE CRITICAL POINT. UCRL-7167, Lawrence Radiation Laboratory, Livermore, California June 1963
148p
Available from CFSTI

Literature values of the vapor pressures of H₂, HD, D₂, N₂, O₂, Ne, Ar, Kr, Xe, CO, NO, CH₄, SF₆, N₂O, CO₂, HCl, HBr, HI, SF₆, H₂S, Cl₂, Br₂, NH₃, C₂H₅OH, SO₂, SO₃, CO₂, HF, HCN, N₂O, CCl₄, and SnCl₂ have been compared with the Krichhoff formula: log Pᵥ = A/T + B, between the normal boiling point and the critical point, where A and B are determined for each liquid. Plots are presented of the deviations computed by the equation: d = Pᵥ - Pᵥₑ, based on accurately determined curves for some of these substances. Empirical hypotheses were developed to allow the prediction of the most probable course of the deviation curve in cases where the data appeared to be unreliable. In addition, where precise values were available, the possibility of the existence of fine structure curvature has been demonstrated. The deviation curves have been used to obtain tables of interpolated vapor pressure values which are, in most cases, most accurate by one order of magnitude than any now in the literature, particularly where smooths regression curves do not discriminate between random experimental errors and systematic deviations of an arbitrarily chosen formula from the real vapor pressure curve. Since the deviation curve is invariably shown to approach the zero ordinate at the critical point with a large negative slope, the correspondence between the critical pressure and temperature is fixed with a high degree of confidence. A particular result of this work is that the critical pressure of NH₃ was shown to be about one atmosphere higher than the presently accepted value. Our values for the critical points of CH₄, C₂H₆, HCN, and Br₂ have also been involved in extending between various authorities in the literature.

#6906009 1969
Eisenberg, D.; Kauzmann, W.
THE STRUCTURE AND PROPERTIES OF WATER
296p

In this book the authors summarize the many experimental observations on water and point to theoretical discussions which effectively correlate the data. The topics covered include the water molecule and forces between water molecules; the thermodynamic properties of steam; the structures of the ices; the thermodynamic, electrical, spectroscopic, and transport properties of the ices and of liquid water; hydrogen bonding in ice and water; and models for liquid water. The authors emphasize the relation of the properties of ice and water to their structures, and discuss properties such as infrared and Raman spectra because they reveal a great deal about the structures. The inclusion of some background material in physical chemistry makes the book accessible to the biologist, biochemist, and geologist as well as to the chemist and physicist. An addendum listing a number of recent articles on the structures and properties of water completes the volume.
This report is a compilation of the physical properties of the metals aluminum, cobalt, iron, magnesium, molybdenum and nickel, and their alloys. The properties included are thermal conductivity, linear thermal expansion, specific heat, electrical resistivity, density, emissivity, diffusivity and magnetic permeability. Data are compiled as a function of temperature, where possible, over the range -457°F to 4500°F. The data contained in this report is primarily derived from data sheets supplied by companies and organizations who responded to an ASTM-ASME survey, trade bulletins, published trade literature and U.S. Government publications.

Eldridge, E. A.; Deen, H. W.
REPORT ON PHYSICAL PROPERTIES OF METALS AND ALLOYS PRISTIC OGENIC TO ELEVATED TEMPERATURES
April 1961
266p

In this work the specific heat values of four refrigerating fluids (R12, R115, R114, and R113) as well as of propane have been measured in the temperature range between 20 and 90°C, for pressures between 0, 2 and 8 absolute atmospheres; the flow-method has been used and a relative accuracy of approximately 0.1% was attained. As a result of the high accuracy of these measurements, the residual dα/Cp which for the maximum pressures used is of the order of 10% of the total Cp value, can be evaluated with a relative accuracy of 1 to 2%. These experimental results, together with the dα/Cp data for water, measured by Vukalovitch et al. and the Cp data for ammonia measured by Osborne et al. have been used to obtain empirically a universal expression for the residual part of the specific heat of gases.

Flann, D. R.; Peavey, B. A., Jr. (Editors)
THERMAL CONDUCTIVITY, PROCEEDINGS OF THE SEVENTH CONFERENCE HELD AT THE NATIONAL BUREAU OF STANDARDS, GAITHERSBURG, MARYLAND, NOVEMBER 13-16, 1967
NBS Special Publication 302, National Bureau of Standards, Washington, D. C.
September 1968
801p

Available from the Superintendent of Documents, GPO.

The present volume contains critical reviews of the literature pertinent to binary equilibrium published through December 1961. The present volume is a supplement to the basic volume Constitution of Binary Alloys by Hansen and Anderko. A total of 1500 system references were consulted in preparing the volume. Written reviews are included for 1719 binary systems, augmented with 435 figures. For the approximately 7-year period since publication of Constitution of Binary Alloys, reviews have been prepared for 753 systems for which previously no data were available (compared to 506 for the 22-year period between the two editions of Hansen). Constitution of Binary Alloys and its first supplement now make written reviews available for 2067 binary systems.

Elliott, J. N.
TABLES OF THE THERMODYNAMIC PROPERTIES OF HEAVY WATER
AECL-1673, Atomic Energy of Canada Ltd., Chalk River, Ontario
January 1963
90p

The Engineering Sciences Data, Chemical Engineering Series contains graphical and tabular data for the thermal conductivity of liquid aliphatic hydrocarbons, liquid alcohols, and water substance; the specific heat at constant pressure of water substance (ice, water and steam, solid, liquid and gaseous heavy water, and sea water); dynamic viscosity and density of water substances (ice, water and steam, liquid and gaseous heavy water, and sea water); and thermal conductivity of liquid halogenated aliphatic hydrocarbons including commercial refrigerants. Each section contains an index of property entries.

Elliott, R. P.
CONSTITUTION OF BINARY ALLOYS, FIRST SUPPLEMENT
1965
877p

The experimental and theoretical calculation of the work functions of particular materials (elements and compounds) are done by various methods: thermionic, photoelectric, field-emission, effusion, contact potential difference, calorimetric and empirical relationships linking the work function to various physical characteristics which are utilized in the theoretical computation of the work function. Richardson constants are also given and references are listed at the end of the book.

Elliott, R. P.
CONSTITUTION OF BINARY ALLOYS, FIRST SUPPLEMENT
1965
877p

154
Commercially available gas, liquid, and chemical sources of oxygen for use with fuel cell batteries are compared. Cryogenic sources are shown to be the most efficient on a weight and volume basis. Chemical fuel generators are satisfactory for applications requiring rather large quantities of oxygen gas at infrequent periods. Compressed gas cylinders are convenient when small quantities of oxygen are desired. A bibliography of selected publications during the past five years is included.

This book contains a compilation of critical solution temperatures (CST), a description of the methods used to determine the CST, a guide to the use of CST data, materials for choosing extraction solvents, and a guide to methods for estimating the CST for untested systems. More than 6000 CST observations are listed. Seventy percent of the systems listed have a hydrocarbon as one component; however, nearly 1100 nonhydrocarbon solvents are listed. In Table I the solvents are listed alphabetically and CST with references are given. Table II gives structure, aniline, and furfural points for high molecular weight hydrocarbons. About 800 aniline points of CST of hydrocarbons with aniline are listed. Table III gives lower critical solution temperatures and upper critical end point or critical temperature, upper layer. Tables IV to VIII present in concise form the data from 5 papers each giving miscibilities of a group of substances. A bibliography of 495 references is included.

This work provides empirical formulae and a nomogram for the calculation of the viscosities of vapors of pure hydrocarbons and of petroleum products. The calculated and experimental data are compared in figures and tables, showing viscosity graphs for hydrocarbons and the dependence of viscosity on molecular weight. References are included.

The literature sources of heat-capacity data on copper, silver, and gold between 0 and 700 K have been compiled and the data critically analyzed. Tables of heat capacity (C_p), enthalpy (H-TS), entropy (S^E), Gibbs energy (G-H^E), and Gibbs energy function (G-H^E)/T have been obtained from the analyses. The literature values of the heat capacity, the electronic coefficient of heat capacity (gamma), and the characteristic temperature [epsilon(n)] are compared with the selected values. The sources of the data are tabulated chronologically along with the temperature range of measurements, purity of sample, and the pertinent experimental procedures used. A bibliography of the references is listed. A brief appraisal of low-temperature calorimetry is given.

This two volume handbook contains tables of thermodynamic properties of 335 gases, 44 liquids and 45 solid substances in all 424 components formed by the following 35 elements and isopettes: H, D, T, N, Ne, Li, B, Be, C, N, O, F, Ne, Na, Mg, Al, Si, P, S, Cl, Ar, X, K, Ca, Br, Kr, Rh, Sr, Zr, Hg, Ph. The thermodynamic properties are calculated for the temperature interval of 293.15 up to 700 000 K. Tables of virial coefficients are given for 34 gases. Volume I of this handbook contains methods of calculation of thermodynamic properties of individual substances. Critical analysis of thermodynamic literature data up to and including 1916 is given on the,constants necessary to calculate tables of thermodynamic properties, such as molecular constants, heat of formation and phase transitions, heat capacity, dissociation energy, etc. The reason for the choice of these constants in the handbook is given as well as the estimates in the cases when experimental data are absent. Volume 2 of the handbook contains tables of thermodynamic properties of the individual substances. All the tables have been calculated by the authors of the handbook on the basis of the data chosen in Volume I.

This annotated bibliography emphasizes phase relationships and their fabrication as evidenced in the physical properties, refractories, and powder metallurgy of molybdenum, niobium and tungsten compounds which are formed with boron, carbon and silicon. The period 1955-1962 yielded the majority of references included in this bibliography.

The literature sources of heat-capacity data on copper, silver, and gold between 0 and 300 K have been compiled and the data critically analyzed. Tables of heat capacity (C_p), enthalpy (H-TS), entropy (S^E), Gibbs energy (G-H^E), and Gibbs energy function (G-H^E)/T have been obtained from the analyses. The literature values of the heat capacity, the electronic coefficient of heat capacity (gamma), and the characteristic temperature [epsilon(n)] are compared with the selected values. The sources of the data are tabulated chronologically along with the temperature range of measurements, purity of sample, and the pertinent experimental procedures used. A bibliography of the references is listed. A brief appraisal of low-temperature calorimetry is given.

This set of books is an evaluated compilation of all data on the thermophysical properties of solid
materials published during the period 1940-1957. Materials with melting points above 1000°F are included. The following physical properties are included: density, melting point, latent heat of fusion, enthalpy, and entropy of solid, liquid, and gaseous argon; total heat of sublimation, specific heat, thermal conductivity, thermal diffusivity, emissivity, reflectivity, linear thermal expansion, vapor pressure and electric resistivity. Volume I contains data for the elements: Volume II, alloys; Volume III, ceramics; Volume IV, cermets, intermetallics, polymeric materials, and composite materials. Volume V, the Appendix, contains the Materials Index, and an alphabetical list of materials. Wherever possible, a recommended value is given.

#6500310 8.133
Goodwin, R. D.; et al.
PRESSURE-DENSITY-TEMPERATURE RELATIONS OF FLUID PARAHYDROGEN FROM 15 TO 100K AT PRESSURES TO 350 ATMOSPHERES
March-April 1962

Experimental data are presented at closely spaced intervals of temperature and density. The range of experimental data is limited to 100 K and to 350 atm less than the critical density. In addition, there are presented uniformly interpolated tables in density and temperature and in pressure and temperature.

#6903054 8.133
Gosman, A. L.; McCarron, R. D.; Hust, J. G.
THERMODYNAMIC PROPERTIES OF ARGON FROM THE TRIPLE POINT TO 300 K AT PRESSURES TO 1000 ATMOSPHERES
March 1969
146p
Available from the Superintendent of Documents, GPO

Tabular values of density, internal energy, enthalpy and entropy of liquid and gaseous argon are presented for temperatures from 83.8 to 300 K at pressures of 0.01 to 1000 atmospheres. Diagns of specific heats, compressibility factor, and entropy are included. The properties presented are calculated from an equation of state which was fitted to experimental P-V-T data from the world literature. Extensive comparisons were made between the equation of state and the experimental data, and deviation plots are presented. The second virial coefficient and Joule-Thomson inversion curve were also calculated and compared.

#6906008 4.183-8.183
Grigsby, D.L.; Welles, S.J.
A BIBLIOGRAPHY ON SELECT PROPERTIES OF ALUMINUM ELECTRONIC (CELL Tin) PROPERTIES
Inromations Center, Hughes Aircraft Company, Culver City, California
April 1969
59p

This bibliography on pure aluminum and dilute aluminum alloys contains references on the following select properties: macroscopic structure, energy band structure, physical and electrical properties, magnetic properties, optical properties, thermal properties, and Debye temperature and attenuation properties.

#6500653 4.133-3.133-8.133
GROUPS IV, V, AND VI TRANSITION METALS, SELECTED REFERENCES ON PREPARATION, STRUCTURE, PHYSICAL PROPERTIES, METALLOGRAPHY, AND TRANSPORT PROPERTIES
ORNL-RMIC-3, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee
April 1965
94p

This bibliography contains lists of selected references on preparation methods, crystal structure, transport properties, and physical and mechanical properties for the Groups IV, V, and VI transition metals. An abstract is included along with each reference entry.

#6902097 4.133-3.133-8.133
GROUPS IV, V, AND VI TRANSITION METALS, SUPPLEMENT NO. 1, SELECTED REFERENCES ON PREPARATION STRUCTURE, PHYSICAL PROPERTIES, METALLOGRAPHY, AND TRANSPORT PROPERTIES
ORNL-RMIC-8, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee
April 1965
27p
Available from CFSTI

This bibliography is a supplement to ORNL-RMIC-3 (Rev.) of April 1965, and contains lists of selected references on preparation methods, crystal structure, transport properties, and physical and mechanical properties for the Groups IV, V, and VI transition metals.

#6500465 8.133-4.133
Gubareff, G.G.; Janssen, J.E.; Torborg, R.H.
THERMAL RADIATION PROPERTIES SURVEY, A REVIEW OF THE LITERATURE SECOND EDITION
Minneapolis, Minnesota: Honeywell Regulator Company 1960
295p

Thermal radiation property data for more than 40 metals and their alloys are given. The properties reported are emissivity, absorptivity, reflectivity, and transmissivity at various surface conditions, temperatures, and wavelengths. In addition, radiation properties of building materials, paints, glasses, paints, and other miscellaneous solid materials have been compiled. Solar absorptivity and equilibrium temperature data for various materials have been tabulated. These values have been gathered from about 200 references in English, German, French, and Russian and have been carefully analyzed to obtain the most trustworthy data for practical application. Recommended data for technically important materials are also presented. An extensive survey was conducted to compile values for Stefan-Boltzmann's, Planck's, and Wien's constants, and the most reliable values for practical application are listed in a separate table.

#6809006 8.133
Harlow, L.
THERMODYNAMIC PROPERTIES OF AMMONIA AS AN IDEAL GAS
August 1968
10p
Available from the Superintendent of Documents, GPO

Thermodynamic functions for ammonia as an ideal gas at one atmosphere pressure have been evaluated. The contribution of the highly anharmonic out-of-plane vibrational mode, including its large coupling with rotation and its coupling with the other vibrational modes, is considered in detail. Tables of C=P/(H - E)/RT, (E-G)/RT, and S/R have been calculated at closely spaced intervals from 50 to 5000 K within an overall uncertainty of less than 0.1 percent at 1000 K.
This monograph contains a consistent set of tables of thermodynamic properties of a large number of diatomic hydrides, deuterides, and tritides, for the ideal gas state at one atmospheric pressure. In addition to the thermodynamic properties of the molecular gases, the tables also include thermodynamic properties for chemical reactions involving the isotopic exchange of hydrogen. The thermodynamic properties tabulated are the heat capacity, enthalpy, Gibb's free energy, and entropy.

This report consists of an index by substance and associated physical and/or thermodynamic property (Substance-Property Index) and a bibliography covering the literature on organic substances for the year 1967. As a further aid to users of this index, a "Guide to the Substance-Property Index" based on the standard order of arrangement of the elements and compounds is included in this report. Work in progress throughout the world dealing with physical, thermochanical, and thermodynamic measurements on organic substances and mixtures is summarized in this index by letter-number references, such as A-67, 4-42, etc. to the abstracts found in the IUPAC Bulletin of Thermodynamics and Thermochemistry, No. 11, June 1968. A new feature this year in the bibliography is the incorporation of the title of each article cited.

A bibliography of 450 references is presented for mechanical, thermal, and transport properties of argon from 0 to 300 K. Each article has been reviewed and coded with regard to properties studied, type of article (i.e., experimental, theoretical, etc.), and method of presentation of data. The temperature and pressure ranges for each property under consideration are also given. An index has been prepared according to property with 4 sub-categories: solid, liquid, gas up to 200 K, and gas above 200 K.

A resume is given of the determinations of the value of the Faraday. Values obtained by silver deposition, iodide oxidation, oxalate oxidation, the proton, and silver diffusion are reviewed. All values are converted to the unified C12 international scale of atomic weights using the international atomic weights of 1967. Values of the Faraday are given in terms of both the NBS (legal) and absolute units of electrical measure. In the latter the new value for the acceleration due to gravity is used in computing the absolute value of electric current. On this basis and using the atomic weight of silver determined by Shields, Craig, and Diebel, and converting to the C12 scale, the value of the Faraday is 96,486.921.6 absolute coulombs per gram-equivalent which differs by only 1 part per million from the value recommended by the National Academy of Sciences-National Research Council. If the atomic weight of silver recommended in 1967 by the International Atomic Weight Commission is used, the Faraday on the new gravity value is 96,486.921.6 absolute coulombs per gram-equivalent which differs by 5 parts per million from that recommended by the National Academy of Sciences-National Research Council. No change in the value of the Faraday adopted by the NAS-NRC Committee is recommended.
Hamers, W. J.

STANDARD CELLS, THEIR CONSTRUCTION, MAINTENANCE, AND CHARACTERISTICS

NBS Monograph 84, National Bureau of Standards, Washington, D. C.

January 15, 1965

42p

Available from the Superintendent of Documents, GPO

This Monograph contains information on the construction, maintenance, and characteristics of standard cells. The effects of temperature, pressure, electric current, light, shock, and vibration on standard cells are discussed. A history of the realization and maintenance of the unit of electromotive force is also included. A record of international comparisons of the unit of electromotive force is presented as well as information on the constancy of the National Reference Group of Standard Cells.

Hamers, W. J.

THEORETICAL MEAN ACTIVITY COEFFICIENTS OF STRONG ELECTROLYTES IN AQUEOUS SOLUTIONS FROM 0 TO 100°C

NSRS-NS-24, National Bureau of Standards, December 1968

271p

Available from the Superintendent of Documents, GPO

In determining the activity coefficients of electrolytes in aqueous solutions from the freezing point to the boiling point of the solvent, various equations have been used in the treatment of the data. This paper gives equations for the activity coefficients of electrolytes of various valence types from 0 to 100°C, and for ion strengths from zero to 0.1 molar or 0.1 molar, as calculated by seven different equations based on the theory of interionic attraction. These equations are those of Debye and Hückel, Glütenberg, Sachtcher, and Sjerrum, and what may be termed an extended Glütenberg equation and an extended Sachtcher equation.

Hanley, H. J. M.

THE VISCOSITY AND THERMAL CONDUCTIVITY COEFFICIENTS OF DILUTE ARGON BETWEEN 100 AND 2000°K


March 1966

Available from the Superintendent of Documents, GPO

The variation of the force constants of the Lennard-Jones, Exp: 6, and Kihara potential functions was investigated by comparing the Chapman-Enskog kinetic theory expression for the viscosity coefficient with the experimental viscosity of dilute argon. It was found that this variation was more pronounced than expected. It was necessary to rationalize the choice of the force constants of each function before using the function to compare theory with experimental data. Of the three, the Kihara was found to give the best correlation, and tables of the viscosity and thermal conductivity coefficients of dilute argon between 100 and 2000°K were computed from this potential and the Chapman-Enskog equations.

Hanley, H. J. M.; Childs, G. E.

DISCREPANCIES BETWEEN VISCOSITY DATA FOR SIMPLE GASES

Science, 159:1114-1117

March 8, 1968

It has been known for some time that Kestin and his co-workers have reported dilute gas viscosity coefficients which differ from the usually accepted values. Recent work from the Los Alamos Scientific Laboratory supplements Kestin's results. It is shown that there is no evidence for not accepting these different data. The whole subject of dilute gas viscosity measurements above room temperature should be reexamined both from the experimenter's and correlator's viewpoints. There is evidence that published tables may be incorrect by as much as 10 percent above 600°K.

Hanley, H. J. M.; Childs, G. E.

THE VISCOSITY AND THERMAL CONDUCTIVITY COEFFICIENTS OF DILUTE NEON, KRYPTON, AND XENON

NBS Technical Note 352, National Bureau of Standards, Washington, D. C.

March 1967

24p

Available from the Superintendent of Documents, GPO

The coefficients of viscosity and thermal conductivity for dilute neon, krypton, and xenon were examined by a method already proved successful for dilute argon, oxygen, and nitrogen. This method selects a suitable potential function, and its parameters, which is then used to correlate theory with experimental data, given the kinetic theory expressions for transport coefficients. The method has recently been expanded and generalized and the results of this general study are applied to this note. The potential functions examined were members of the m-6, Kihara, Exp: 6 and Morse families. It was found that the Kihara was most suitable for neon, and the m-6, with m = 17 and m = 24, was most suitable for krypton and xenon, respectively. Viscosity and thermal conductivities were calculated from these functions and tables are given between 100 and 1000°K.

Hanley, H. J. M.; Klein, M.

ON THE SELECTION OF THE INTERMOLECULAR POTENTIAL FUNCTION: APPLICATION OF STATISTICAL MECHANICAL THEORY TO EXPERIMENTAL DATA

NBS Technical Note 360, National Bureau of Standards, Boulder, Colorado

November 20, 1967

88p

Available from the Superintendent of Documents, GPO

A method has been developed to evaluate quantitatively the relationship between model intermolecular potential functions and macroscopic experimental properties. Specifically, the following function families have been studied: m-6, Kihara, Exp: 6, and Morse; the properties studied are: viscosity coefficient, diffusion coefficient, second virial and Joule-Thomson coefficients. The principal conclusions from this work are: 1. a temperature range exists for each m that is insensitive to the potential function; 2. the function families studied are essentially equivalent; 3. more than one function is required to represent data over a range of about 1000°K; 4. the experimental error which can be tolerated if experimental data are to be used to select a potential function has been estimated. The conclusions are experimentally verified for both transport and equilibrium properties.

Hanley, M.; Anderko, K.

CONSTITUTION OF BINARY ALLOYS, SECOND EDITION


305p

This monograph deals with the composition and crystal structure of 1,334 binary alloy systems, compiled from a critical review of the literature. The composition data are presented in the form of phase diagrams for those systems for which such diagrams had been published. Other systems have information about mutual solubility and melting points compiled. The crystal structure data includes symmoric class and lattice parameters. An index to systems is provided.
Basic theory of electrolytic solutions is presented in the first five chapters of this book. Chapters 2, 3, and 4 present interionic attraction theory, properties of ionic atmosphere, the thermodynamic properties of aqueous solutions and the theory of irreversible processes in electrolytic solutions. Numerical compilations of physical constants, characteristic ionic and mathematical functions are presented in chapter 5. The experimental methods discussed in chapters 6 through 10 are irreversible processes in solutions of strong electrolytes; conductance, transference numbers, viscosity, diffusion; coulomb forces and ion association, weak electrolytes, frequency and field effects; thermochemical quantities, including partial molal volumes, coefficients of expansion and compressibility; the calculation of activity and osmotic coefficients from freezing points, boiling points and vapor pressures; and the thermodynamics of galvanic cells. No apparatus or experimental techniques are discussed in detail. In the last third of the book, the emphasis is upon the properties of electrolytes rather than on the methods by which they were obtained. Chapters 11, 12 and 13 deal with solutions of hydrochloric acid, 1-1 electrolytes and polyvalent electrolytes, respectively. Mixtures of strong electrolytes in water, the sublimation of solid electrolytes, the ionic constants of weak electrolytes, their temperature coefficients and the ionization of weak electrolytes in salt solutions are considered in chapters 15. Extensive tables of quantities derived from the best available data are given in the appendix. Author, subject and chemical indexes are included.

Harned, H. S.; Owen, B. B.
THE PHYSICAL CHEMISTRY OF ELECTROLYTIC SOLUTIONS, THIRD EDITION
New York: Reinhold Publishing Corporation
1958
803p

#6007009

Hilsenrath, J.; Klein, M.
TABLES OF THERMODYNAMIC PROPERTIES AND CHEMICAL COMPOSITION OF NITROGEN IN CHEMICAL EQUILIBRIUM INCLUDING SECOND VIRIAL CORRECTIONS FROM 1600 K TO 15,000 K
April 1966
307p
Available from DDC

Tables for the thermodynamic properties for nitrogen are presented which take into account the effect of dissociation and ionization and the limiting-law Debye-Hückel and second virial corrections, upon the thermodynamic properties and the chemical equilibrium compositions. Values are from 1600 K in steps of 100 K to 15,000 K at close spacings in the logarithm of the density \[ \log \rho / \rho_0 = -7.0(0.2,2.2) \] for the compressibility factor \( Z = \rho V / RT \); the dimensionless functions for: internal energy, \( E/RT \); enthalpy, \( H/RT \); entropy, \( S/RT \); \( \log P(\text{atm}) \) and \( Z^* = Z \); and the mole fractions of the species in chemical equilibrium. The underlying equations and the input data are discussed briefly. The effects of the real gas corrections on the equilibrium properties are illustrated graphically. The equilibrium composition is given for selected temperatures over the tabulated density range. The wide range of temperatures and densities over which the thermodynamic properties have been tabulated make the tables useful in a variety of engineering design and test programs, and in scientific research and development.

Hilsenrath, J.; Klein, M.
TABLES OF THERMODYNAMIC PROPERTIES OF AIR IN CHEMICAL EQUILIBRIUM INCLUDING SECOND VIRIAL CORRECTIONS FROM 1500°K TO 15,000°K
AEDC-TR-65-58, Arnold Engineering Development Center, Arnold Air Force Station, Tennessee
March 1965
533p
Available from DDC

Tables for the thermodynamic properties for air are presented which take into account the effect of dissociation and ionization and the limiting-law Debye-Hückel and second virial corrections upon the thermodynamic properties and the chemical equilibrium compositions. Values are from 1500°K in steps of 100°K to 15,000°K at close spacings in the logarithm of the density \[ \log \rho / \rho_0 = -7.0(0.2,2.2) \] for the compressibility factor \( Z = \rho V / RT \); the dimensionless functions for: internal energy, \( E/RT \); enthalpy, \( H/RT \); entropy, \( S/RT \); \( \log P(\text{atm}) \) and \( Z^* = Z \); and the mole fractions of the species in chemical equilibrium. The underlying equations and the input data are discussed briefly. The effects of the real gas corrections on the equilibrium properties are illustrated graphically. The equilibrium composition is given for selected temperatures over the tabulated density range. The wide range of temperatures and densities over which the thermodynamic properties have been tabulated make the tables useful in a variety of engineering design and test programs, and in scientific research and development.

Hilsenrath, J.; Klein, M.
TABLES OF THERMODYNAMIC PROPERTIES OF NITROGEN IN CHEMICAL EQUILIBRIUM INCLUDING SECOND VIRIAL CORRECTIONS FROM 2000°K TO 15,000°K
AEDC-TRD-63-162, Arnold Engineering Development Center, Air Force Systems Command
March 1964
162p
Available from DDC

Tables for the thermodynamic properties for nitrogen are presented which take into account the effect of dissociation and ionization and the
limiting-law Debye-Hückel and second virial corrections, upon the thermodynamic properties and the equilibrium compositions. Values are tabulated from 2000°K in steps of 100°K to 15,000°K at close spacings in the logarithm of the density \( \log \rho \). The underlying equations and the input data are discussed briefly. The effects of the real gas corrections on the equilibrium properties are illustrated graphically. The equilibrium composition is given for selected temperatures over the tabulated density range. The wide range of temperatures and densities over which the thermodynamic properties have been tabulated make the tables useful in a variety of engineering design and test programs, and in scientific research and development.

#6905007
Hilsenrath, J.; Messina, C. G.; Evans, W. H.
TABLES OF IDEAL GAS THERMODYNAMIC FUNCTIONS FOR 73 ATOMS AND THEIR FIRST AND SECOND IONS TO 10,000°K
Technical Document AFWL-TR-64-44, Air Force Weapons Laboratory, Kirtland Air Force Base, New Mexico
August 1964
44p
Available from DDC

Tables are presented for five thermodynamic properties, \( C_{p} \), \( C_{v} \), \( H^{m} \), \( S^{m} \), and \( Y_{\text{ideal}} \), and the thermodynamic property for the ideal gas. The tables are given at 100° intervals starting at 100°K for the atoms and at 100° or 1000° for the ions and extending to 10,000°K. They were computed for the atoms and atomic ions considered as ideal monatomic gases by direct summation over the available energy levels. The summation over energy levels was made from the ground state up to the principal quantum number \( n \). Where the quantity \( n \) for the ground state exceeds 5, summations were made only over levels having the same \( n \) as the ground state. The sensitivity of the thermal functions to the inclusion of higher energy states in the partition functions is discussed briefly, graphically for the specific heats, and tabulated for the Gibbs free energy function.

#6900343
Hilsenrath, J.; Ziegler, G. G.
EINSTEIN FUNCTIONS
NBS Monograph 49, National Bureau of Standards, Washington, D.C.
July 1962
258p
Available from the Superintendent of Documents, GPO

Tables are presented for the contribution of a harmonic oscillator to the free energy function, enthalpy function, entropy and heat capacity of gases. Dimensionless values of the Planck-Einstein functions are given as a function of the temperature \( T \) and \( \log h_{c} \). A second table gives the contributions in calories/mole \( \text{cal/mole} \) as a function of frequency \( \nu \). A second table gives the contributions in calories/mole \( \text{cal/mole} \) as a function of frequency \( \nu \). The temperature \( T \) was computed using the values 1.43380 for the second radiation constant \( c_{r} \), and 1.98177 for the universal gas constant \( R \). The arguments for the latter table are spaced at 10 wave number intervals from 1000 cm\(^{-1}\) to 4000 cm\(^{-1}\).
This technical report covers work in a continuing systematic program on the thermophysical properties of materials involving the literature search, collection, codification, organization, data evaluation, correlation, analysis, and synthesis, the preparation of "intermediate tables" presenting the total available experimental information, and the final preparation of internally consistent tables of "best data" referred to as "Tables of Recommended Most Probable Values." The work reported on consists of data tables projects on the thermal conductivity, specific heat, thermal radiant properties (emittance, reflectance, absorptance, transmittance), thermal diffusivity, and thermal expansion of elements, alloys, compounds, mixtures, and systems, and the thermal conductivity, specific heat and viscosity of liquids and gases. Property data are presented in both tabular and graphical forms, with accompanying tables giving specifications and characterizations of the test specimens. The resulting data tables are disseminated at large through the 13-volume TPRC Series on Thermophysical Properties of Matter, while the resulting bibliographic information is disseminated through the Thermophysical Properties Research Literature Retrieval Guide. This report does not contain the completed thousands of data sheets, but does contain the Appendix, the Table of Contents and the Grouping of Materials and List of Figures and Tables for each of the first seven volumes (which contain over 8000 pages) of the TPRC Series to show the scope of their coverage.

Ho, C. Y.; Powell, R. W.; Liley, P. E.
THERMAL CONDUCTIVITY OF SELECTED MATERIALS, PART 2
NSRDS-NBS-10, National Bureau of Standards, Washington, D. C.
February 1968
146p
Available from the Superintendent of Documents, GPO

The work presented in this report comprises the critical evaluation, analysis, synthesis of the available thermal conductivity data, and the generation of recommended values for twelve metallic elements, mainly for the solid state, for a range of graphite and for three fluids in the gaseous state. These are cadmium, chromium, lead, magnesium, molybdenum, nickel, niobium, tantalum, tin, titanium, zinc, zirconium, Acheson graphite, ATJ graphite, pyrolytic graphite, 875S graphite, 9805 graphite acetone, ammonia and methane. For each of the materials recommended values are given over a wide range of temperature.

Ho, J. C.; O'Neal, H. R.; Phillips, N. E.
LOW TEMPERATURE HEAT CAPACITIES OF CONSTANTAN AND MANGANIN
The Review of Scientific Instruments, 34:782-783
July 1963

The heat capacities of Constantan and manganin have been measured from 0.15 and 0.2K, respectively, to 4.2K. Both alloys have large hyperfine heat capacities at a few tenths of a degree, and data are presented which can be used in making corrections to calorimetric measurements in which one of these materials is used as a heater.

Holley, C. (Compiler and Editor)
THERMODYNAMIC AND TRANSPORT PROPERTIES OF URANIUM DIOXIDE AND RELATED PHASES
1965
105p

This report deals with the physical, thermodynamic, and transport properties of uranium dioxide and related phases of the uranium-oxygen system. It includes data and a summary of research work done in the field through 1964. The following topics are included: crystal structure, heat capacity, free energy, enthalpy and entropy measurements, vaporization processes, surface and oxidation properties, thermal conductivity, electrical properties, optical properties, and practical implications of thermodynamic and transport properties.

Horsley, L. H.
AZEOTROPIC DATA
Advances in Chemistry Series 6, American Chemical Society, Washington, D. C.
June 1952
535p

Information is given in tabular form on 14,501 binary systems of azeotropes and nonazeotropes. These include boiling points of the pure constituents and of the azeotropes, and the compounds of the azeotropes when available, are given. The boiling points of the azeotropes may range from less than -100°C to greater than 2300°C. Similar data are given for 407 ternary systems. Included are a formula index, a 437 reference bibliography, and three articles, "Vapor-Liquid Equilibrium Diagrams of Alcohol- Ketone Azeotropes as a Function of Pressure," "Graphical Method for Predicting Effect of Pressure on Azeotropic Systems," and "Graphical Method for Predicting Azeotropism and Effect of Pressure on Azeotropic Constants."

Horsley, L. H.; Taplin, W. S.
AZEOTROPIC DATA II
Advances in Chemistry Series 35, American Chemical Society, Washington, D. C.
1962
105p

This volume is a supplement to AZEOTROPIC DATA published as Advances in Chemistry Series No. 6, American Chemical Society, 1952. It includes revised data on systems in the original table plus new data on azeotropes, nonazeotropes, and vapor-liquid equilibria collected since 1952. No attempt has been made to evaluate the data. Where appreciable differences occur in data from different sources, more than one set of data is recorded. A brief description is included for the calculation of azeotropic data for immiscible systems from vapor pressure data. The tables are arranged in the same manner as the previous volume.

Hughes, E. E.; Liss, S. C.
VAPOR PRESSURES OF ORGANIC COMPOUNDS IN THE RANGE BELOW ONE MILLIMETER OF MERCURY
NBS Technical Note 70, National Bureau of Standards, Washington, D. C.
October 1960
24p
Available from CFSTI, PB 161 571

Data for vapor pressures less than one millimeter of mercury for over three hundred organic compounds have been collected. The values are presented in tabular form. No attempt has been made to evaluate the reliability of these data except where several investigators have studied the same compound over
the same pressure range. A brief discussion of the methods and instruments which have been used to measure low vapor pressures is included.

Hughes, R. L.; Smith, J. C.; Lawless, R. A.
Holzmann, R. T. (Editor)
PRODUCTION OF THE BORANES AND RELATED RESEARCH
1967
533p
This is a review and analysis of borane chemistry. The topics covered include: precursors to the boranes; formation of the boron-hydrogen bond and diborane; preparation and properties of the boron hydrides, alanes, and carboranes; kinetics and mechanism studies; preparation and properties of pentaborane and decaborane from diborane; separation and purification; analytical methods; boron-nitrogen chemistry; and base chemistry and appendices on toxicology; infrared spectra; molecular structures; physical properties of boranes and their derivatives.

**#6511007**  *8.133-4.133*
Hultzgen, R.; Orr, R.L.; Kelley, K.K.
SUPPLEMENT TO SELECTED VALUES OF THERMODYNAMIC PROPERTIES OF METALS AND ALLOYS
California University, College of Engineering and Lawrence Radiation Laboratory, Berkeley, California
June, 1970
505+ loose-leaf pages
This Supplement contains the results of evaluations for 56 metallic elements and 178 binary alloy systems for which thermodynamic data were available. Reevaluation and extension of the selected values will be a continuing process as new data become available. Prior to the publication of a revised edition of the book, all additional evaluations, as they are completed, will be issued in loose-leaf form at frequent intervals for inclusion with this Supplement. Description of the methods of evaluation and forms of presenting the data may be found in the parent volume of "Selected Values." Results of the evaluations in the form of discussions and analyses of the data and tables of selected values are given. Some of the properties covered include freezing point, vapor pressure, transition temperature, phase diagrams, entropy, heat capacity, internal energy, function, heat content; heat of transition, sublimation, and conversion, and vaporization; entropy of transition, sublimation, and conversion, and vaporization; free energy of sublimation and vaporization; free energy of formation; free energy excess integral, relative partial molar and relative partial molar excess; heat of formation - relative partial molar; entropy of formation - relative partial molar and relative partial molar excess; heat capacity - relative partial molar.

**#6500306**  *8.233*
Hust, J. G.; Cosman, A. L.
FUNCTIONS FOR THE CALCULATION OF ENTROPY, ENTHALPY, AND INTERNAL ENERGY FOR REAL FLUIDS USING EQUATIONS OF STATE AND SPECIFIC HEATS
1963
1p
659
Available from the Cryogenic Data Center
General integral equations for the determination of entropy, enthalpy, and internal energy are derived. In the first part of this paper the integration path is taken to be on the real gas surface thus yielding one set of equations. In the latter part of this paper the integration path is defined such that ideal gas specific heats and reference properties are introduced. The latter set of integral equations are applied to a particular equation of state resulting in algebraic expressions for the entropy, enthalpy, and internal energy.

**#6500403**  *8.233*
Hust, J. G.; Stewart, R. B.
THERMODYNAMIC PROPERTY VALUES FOR GASEOUS AND LIQUID CARBON MONOXIDE FROM 70 TO 300K WITH PRESSURES TO 500 ATMOSPHERES
NBS Technical Note 202, National Bureau of Standards, Boulder, Colorado
November 1963
19p
Available from the Superintendent of Documents, GPO
The internal energy, entropy, enthalpy, and density of carbon monoxide are tabulated as functions of pressure and temperature from 70 to 300K and 0.1 to 300 atmospheres. A compressibility factor-pressure chart and a temperature-entropy chart are also included. These P-p-T values were calculated by using the Su principle of corresponding states with nitrogen as a model. Extensive comparisons are included, illustrating the deviations of the calculated values from the experimental and other correlated data. Equations representing the P-p-T surface and the vapor pressure are given.

**#6500482**  *8.133*
Hust, J. C.; et al.
BIBLIOGRAPHY OF THE THERMOPHYSICAL PROPERTIES OF OXYGEN AT LOW TEMPERATURES
NBS Technical Note 137, National Bureau of Standards, Washington, D. C.
February 1962
5p
709
Available from the CPSTI, PR 161 638
This bibliography of the mechanical, thermodynamic, and transport properties of oxygen below 0°C presents 325 references and is the result of a thorough search of the world's scientific and engineering literature. In addition to searching abstracting journals and bibliographies, the authors reviewed each document for property data and for additional references. Listed for each reference are the properties and the corresponding temperature and pressure range, together with additional pertinent information such as the type of data (i.e., whether the data are derived from experimental measurements, theoretical considerations or as a compilation from other sources), the form and amount of data, etc. An index according to property, sub-indexed for temperature and pressure ranges, and an author index is included.

**#6500463**  *8.143-4.143*
Janssen, J.; Luck, J.; Torborg, R.
REFLECTANCE OF ANODIZED TITANIUM AND BERYLLIUM Aeronautical Systems Division, Wright-Patterson Air Force Base, Ohio
(NO date)
5p
738
Selection of materials having the desired thermal radiation properties is of primary importance for space applications, since heat transfer to and from an object in space occurs only by radiation. The effect of anodizing variables on the reflectance of anodized titanium and beryllium was studied over the range of 0.4 to 22 microns in a vacuum of 10^-8 mm Hg with specimen temperatures of 100 to 1300°F. Of the anodizing variables studied, the various anodizing solutions generally had the greatest effect on the reflectance. A number of surfaces investigated had high alpha/epsilon ratios, indicating usefulness as potential solar collector surfaces.
Selection of materials having proper thermal radiation properties permits the engineer to achieve passive temperature control in spacecraft. The effect of anodizing variables on the reflectance of anodized aluminum and magnesium was studied over the range of 0.4 to 22 micron in a vacuum of 10^-5 mm of Hg with specimen temperatures of 100 to 825 F. In general, the anodizing process had a greater effect on the reflectance than did alloying elements in the metal. The pressure of water in some coatings, which was driven off at elevated temperatures in the vacuum, gave lower reflectance in the infrared.

The data in the handbook was taken from results published in the literature; for the most part, numerical values are those in the original source material. The following categories of data are included: physical properties: atomic and ionic radii, melting and boiling points, density viscosity, vapor pressure, surface tension, refractive index, critical data; 2. thermodynamic properties: liquid-liquid immiscibility, metal-molten salt systems, metal oxide-molten salt systems, silver salts-molten salt systems, phase-rule studies, solubility, thermal data, cryoscopy, expansivity, compressibility, and high pressure studies; 3. electrochemical properties: reference electrodes and emf series, electrical conductance, polarography, electromigration, dielectric and thermoelectric properties; 4. spectroscopy and structure: vibrational spectroscopy, visible and U.V. spectroscopy, NMR and EPR spectroscopy, X-ray and neutron diffraction, and fused salt electrolytes; 5. practical features: melt preparation and purification, containment and corrosion, chromatographic analysis, electrolysis, chemical processes, fuel cells, batteries and nuclear technology.

An updating in practical methods for computing thermodynamic properties and a compilation of data and correlations for thermodynamic calculations are presented. Parts of the text deals with the calculation of free energy changes and their use as a criterion of thermodynamic properties from molecular parameters, spectroscopic data, and statistical thermodynamics; and estimation methods. Part II presents 78 tables of data and correlations of the many contributors in the field of estimation of thermodynamic properties. These tables have been collected to make application of the estimation methods as convenient as possible. The tables are reproduced as compiled in the original work of each investigator.

Data on the electrical conductance, density and viscosity of single-salt melts were compiled from a comprehensive search of the literature up to December 1966 and a critical assessment made of the compiled data. Recommended values were determined and are presented as functions of temperature in the form of equations and tables. The results for some 174 compounds as single-salt melts are reported; no attempt was made in the present effort to embrace the results for molten salt mixtures. Data are presented for fluorides, chlorides, bromides, iodides, carbonates, nitrates, oxides, sulfides, sulfates and a miscellaneous group.

This book consists of two sections as follows: Section 1. The critical evaluation of excess free energies of binary molten salt mixtures with a common ion from equilibrium-type electrochemical cells is described in this report. For this purpose calculations using the original emf data were worked systematically undertaken to establish comparisons of free energy values of various workers that would be significant. The reversibility of electrodes is investigated by comparing the electro motive force of cells with a single molten salt as liquid electrolyte with thermochromic data. Section 2. Data on the surface tensions of single salt melts have been systematically collected and evaluated. Results are given for 106 inorganic compounds over a range of temperatures where available.

This is an unevulated compilation of the pKa values for inorganic acids, phosphates, carboxylic acids, phenols, alcohols and oxygen acids, amino acids, peptides, nitrogen compounds, chlor, common acids, and indicators. References are given for each value from a total of 96 sources.

Available from the Superintendent of Documents, GPO

Available from Superintendent of Documents, GPO
This first phase of the Compendium covers ten properties of ten fluids (Part I), three properties of solids (Part II), and an extensive bibliography of references including a Property and Material Index (Part III). In Part I, density, expansivity, thermal conductivity, specific heat and enthalpy, transition heats, phase equilibria, dielectric constants, adsorption, surface tension and viscosity for the solid, liquid and gas phases of helium, hydrogen, neon, nitrogen, oxygen, air, carbon monoxide, fluorine, argon and methane are given wherever adequate data could be collected. In Part II, thermal expansion, thermal conductivity, and specific heat and enthalpy are given for a number of solids of interest in cryogenic engineering. Data sheets, primarily in graphic form, are presented for "best values" of data obtained. The purpose of the thermometric data sheets is to furnish references and tables of selected values with appropriate comments are furnished with each data sheet to document the data presented. Conversion tables and other helpful information are also included.

Keenan, J. H.; Keyes, F. C.
THERMODYNAMIC PROPERTIES OF STEAM including DATA FOR THE LIQUID AND SOLID PHASES, FIRST EDITION
New York: John Wiley & Sons, Inc.
163
89p

The present tables are computed from entirely new formulations of the properties of water. Table 1 gives absolute pressure, specific volume, enthalpy and entropy for saturation temperatures. Table 2 gives specific volume, enthalpy, entropy and internal energy for saturation pressures. In Table 3 specific volume, enthalpy and entropy are given for superheated vapor. Within the limitations of the size of the volume, tables are included giving the viscosity, the heat conductivity, compressed liquid properties, data on ice and its vapor, isentropic expansion exponents, and certain thermometric calibration data, while at the same time retaining the temperature-scale conversion table, logarithm tables, and conversion factor tables of the 1030 volume. Certain charts of the specific heat hitherto not available are included and show the extraordinarily rapid changes and the large values of the quantity corresponding to volumes at and near the critical state. The range of temperature (1600°F) and of pressure (5500 psi) of these tables is greater than the 1930 tables. The inclusion of data for the liquid phase is an innovation made possible by the recent exact measurements of heat quantities and specific volumes. The sources of specific volume data are briefly discussed with references to the devices of formulation employed and the analytical expressions found to be servicable.

Keenan, J. H.; et al.
STEAM TABLES: THERMODYNAMIC PROPERTIES OF WATER INCLUDING VAPOR, LIQUID, AND SOLID PHASES (ENGLISH UNITS)
New York: John Wiley & Sons, Inc.
166
174p

The present Steam Tables are based on the tables of Keenan and Keyes (1956). Included are tables giving the mass, heat, and latent heat of the liquid and of the vapor, enthalpy, heat capacity, P-V-T relation, Gibbs free energy. Also values for the compressed liquid region are inserted. The tables are in good accord with heats at high International Skeleton Tables (1963). The present tables extend the range of the Keenan and Keyes tables to 2400°F and to 15,000 psi. A table of the phase changes accompanying polymorphic transformations is provided, along with a separate and detailed table for states near the critical point. Transport properties, i.e., viscosity and thermal conductivity, are not included.
The first Federal Bureau of Mines compilation of entropy and low-temperature heat-capacity data was Bulletin 350, which appeared in 1932. Revisions and elaborations were issued as Bulletin 394 (1936), Bulletin 434 (1941), and Bulletin 477 (1950). The present bulletin covers data available through September 1959 and is a further elaboration and revision, made desirable by the fact that nearly 1300 entropy values for inorganic substances at 298.15°K now are known, as compared with 800 in the 1950 publication, which covered data available through October 1948. The present bulletin contains the currently pertinent explanatory matter that appeared in its predecessors. It has the same dual purpose of assembling the available entropy values at 298.15°K (25.00°C) for the elements and inorganic compounds and giving enough explanation of methods of measuring and calculating entropies to make the results comprehensible. It should be emphasized that this bulletin supersedes those mentioned above.

This treatise is a source document containing information on helium resources, production, conservation, thermodynamic properties, liquefaction and refrigeration techniques, transportation and storage of liquid and safety requirements. It also contains a discussion of uses for liquid and cold gas in cryoelectronics, superconductivity, bubble chambers, cryopumping and microwave and space systems. The book brings together articles by noted authorities in cryogenic technology in which they discuss their specialized field in great depth.

This textbook deals with theoretical and experimental thermodynamics and thermochemistry and contains tables of thermochemical constants for a large number of elements and compounds of metallurgical importance. Theoretical topics considered include the mass action law, thermodynamic functions and solutions. Experimental methods discussed include calorimetric methods, electrolytic force methods, and equilibria with a gaseous phase and a condensed phase. The estimation of thermochemical data and examples of thermochemical treatments of metallurgical problems are also discussed. The thermochemical data have been compiled from published experimental results of over a century. The authors have, wherever possible, listed the most accurate values and assigned a range of uncertainty to each value. The data have been included for heats of formation, standard entropies, heats of transformation, fusion and evaporation, heat capacities, vapor pressure, standard free energies of reaction, and crystal structure type.

Thermochemical data are given for intermetallic systems and for the transition metals as carbides, phosphides, nitrides, sulphides and oxides. Some of the properties covered include partial free energy, partial heat of solution, partial entropy in solution, integral free energy, integral heat of solution, integral entropy of solution, latent heat of fusion, entropy of fusion, latent heat of transformation, and volume change on formation. References and an index of alloy systems are given at the end to simplify searching.
This section of the Landolt-Bornstein tables contains data on fusion equilibria and interfacial tensions. The section includes data from literature published through September 1955. Where several values for one measurement were found, the average was presented. Data on fusion equilibria are presented for binary and ternary systems of inorganic compounds, alkali metal systems, and organic-inorganic systems. Interfacial tension data are presented for pure liquids towards their vapors or air, solutions towards air, two non-miscible systems, and interfacial films on water. Data are also presented on absorption from the gas phase and the liquid phase, and on paper chromatography.

This section of the Landolt-Bornstein tables contains data on the thermodynamic state functions of gaseous substances in liquids, solid and liquid metals and alloys, as well as the solubilities of solid and liquid substances in liquid solution media. Except for the section "gases in metals," only solution equilibria in the temperature region of about -200°C to 300°C are presented. The data were taken from literature published through 1962. Solution equilibria are presented for gases with water, aqueous solutions, non-aqueous inorganic liquids, and mixtures of organic and inorganic liquids. Also, solution equilibria of hydrogen, oxygen and nitrogen in pure metals are presented. Solution equilibria are presented for inorganic materials in water and other inorganic liquids, organic liquids, organic materials in water, and solutions with organic solvents.

This section of the Landolt-Bornstein tables includes data on solution equilibria of multi-component systems for solutions of solid and liquid organic materials in organic liquids and equilibria in systems with several non-miscible phases.
This section of the Landolt-Bornstein tables contains data on the electrical properties of 
electrochemical systems. The data were taken from 
the literature published through October 1960. The 
data compiled includes the electrical conductivity 
of molten salts, pure liquid, and aqueous 
electrolyte and field solutions; frequency and 
field strength dependence of conductivity of 
aqueous solutions; electrical conductivity of 
non-aqueous solutions; electrophoretic potentials 
and electrokinetic potentials, standard electrode 
potentials, dissociation constants of inorganic 
and organic compounds and acid-base indicators.
systems, units and conversion tables, atomic weights, reduction to standard states, densities of water and mercury. Physical and mechanical properties of non-metallic solids are given for natural and synthetic building materials, organic natural materials (wood, paper, cellulose, wood pulp), fibrous materials, ceramics, glasses, synthetic materials, and natural synthetic materials. Also presented are data on friction and rolling resistance of non-metallic and metallic substances, viscosity, flow through tubes, circulatory flow of bodies during two dimensional flow and rough flow, flotation, deep offon, sound, radiation and absorption, the human ear, sound spectra, vibrational numbers, spatial and building acoustics, sound recording and biological effects of ultrasounics.

#6500539
LANDBOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL
RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY,
PHYSICS AND TECHNOLOGY, SIXTH EDITION,
VOLUME IV: TECHNOLOGY,
PART 2a: MATERIAL VALUES AND PROPERTIES OF
INDUSTRIAL METALS, BASIC PRINCIPLES, TESTING
PROCESSES, INDUSTRIAL STEELS
Borchers, H.; Schmidt, E. (Editors)
Berlin, New York: Springer-Verlag
1967
889p
In German

This section of the Landolt-Bornstein tables includes data on metallographic principles and concepts, testing properties, and the properties of iron and its alloys. Data are given on diagrams of static and dynamic and plastic deformation of steel materials. The physical properties of iron tabulated are lattice constants, density, thermal expansion, diffusion, atomic heats, specific heats, enthalpies, or entropies and free enthalpies, phase diagrams and transition points, thermal conductivity, electrical properties, magnetic properties, optical constants and excitation cross-sections. Composition, treatment, strength values and mechanical and chemical properties of pure iron, unalloyed steels, highly alloyed steels, special steels, and cast iron are presented.

#6804008
LANDBOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL
RELATIONSHIPS IN PHYSICS, CHEMISTRY, ASTRONOMY,
PHYSICS AND TECHNOLOGY, SIXTH EDITION,
VOLUME IV: TECHNOLOGY,
PART 4a: HEAT TECHNOLOGY, HEAT TECHNOLOGY
ENGINEERING MEASURING METHODS, THERMODYNAMIC
PROPERTIES OF HOMOGENEOUS MATERIALS
Hausen, H. (Editor)
Berlin, New York: Springer-Verlag
1967
944p
In German

This section of Landolt-Bornstein is a compilation of data dealing with the technology of thermal measurements and the thermodynamic properties of gases, vapors, liquids, and solid materials. Data are included for such areas as the basis of temperature measurement, the determination of temperature measurement with contact thermometers; optical temperature measurement (Pyrometry); hygrometry; thermodynamic properties of homogeneous materials, gases, water and steam, refrigerating agents; thermal expansion of solids and liquids; and the specific thermal properties of technically important materials.

#6802001
LANDBOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL
RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW
SERIES,
GROUP II: ATOMIC AND MOLECULAR PHYSICS,
VOLUME 5: MOLECULAR ACOUSTICS
Schaaffs, W.
Heiwege, K.-H.; Heiwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1967
280p
In German and English

This volume lists data on acoustics based essentially on the molecular properties of matter, and values of sound velocity, absorption coefficients, and dispersion data. The measurements were made at frequencies ranging from 10 Kc to 200 Mc. Preference was generally given in compiling data to measurements for which statements on density, chemical purity, and the temperature coefficients were added. Often, however, several statements can be found for a measured value. Generally, the limit of accuracy is 1% to 2%. Data are given on sound velocity in gases and liquids; for pure gases and liquids as a function of temperature, pressure, and structure; also for mixtures of gases, mixtures of liquids, and solution. Data are compiled on sound absorption and dispersion in gases; sound absorption in homogeneous liquids, in solution, and on sound propagation in polymers, also on sound propagation in isotropic or quasi-isotropic solids (metals, inorganic, or non-metallic, organic solids). There are special sections on sound propagation in liquid helium and shock wave velocity.

#6608030
LANDBOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL
RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW
SERIES,
GROUP III: CRYSTAL AND SOLID STATE PHYSICS,
VOLUME 1: ELASTIC, PIEZOELECTRIC, PIEZOPTIC AND
ELECTROTROPIC CONSTANTS OF CRYSTALS
Bechmann, R.; Heiwege, R.F.S.
Heiwege, K.-H.; Heiwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1966
160p
In German and English

This volume of the new Landolt-Bornstein series gives the elastic, piezoelectric, piezooptic, and electrooptic constants and other data included were taken from data published in the literature through 1965. For materials where one or two sets of measurements of a given constant were found, those quoted as published; for more than two sets of measurements, the mean and coefficient of variation are given. For non-piezoelectric constants, the elastic stiffness and compliance constants, together with their temperature and pressure variations, are given. For piezoelectric constants, the elastic, piezoelectric, dielectric, piezooptic, and electrooptic constants, plus the electromechanical coupling factors are given. The temperature coefficients of the elastic stiffness, compliance, piezoelectric stress and strain constants are also given.

#7001036
LANDBOLT-BORNSTEIN, NUMERICAL DATA AND FUNCTIONAL
RELATIONSHIPS IN SCIENCE AND TECHNOLOGY, NEW
SERIES,
GROUP III: CRYSTAL AND SOLID STATE PHYSICS,
VOLUME 2: ELASTIC, PIEZOELECTRIC, PIEZOPTIC,
ELECTROTROPIC CONSTANTS AND NONLINEAR DIELECTRIC
SUSCEPTIBILITIES OF CRYSTALS
Heiwege, K.-H.; Heiwege, A.M. (Editors)
Berlin, New York: Springer-Verlag
1969
242p
In German and English
Volume III/2 of the New Series of Landolt-Bornstein continues Volume III/1 only three years after publication of the latter. All the tables contained in III/1 are brought up to date by supplements in III/2. In addition, the elastic and electrooptic constants of higher order and the nonlinear optical susceptibilities have now been included. Readers should note that the Index of Substances at the end of III/2 refers to both volumes.

#6911030 8.153
Landshoff, R. K. M.; Nagel, J. L. (Editors)
THERMAL RADIATION PHENOMENA VOLUME I RADIATIVE PROPERTIES OF AIR
New York: IFI/Plenum 1969 664p

The editors of this book have prepared an introduction for the tables presented in DASA 1917-3. A few more figures have been added and a number of errors have been corrected. This volume presents tables of the equilibrium radiative properties and its components for a wide range of temperatures and densities. The work is divided into five parts: Part A is a summary section designed to give an overview of the field. Some of the data are presented in semi-schematic form and should not be used for quantitative work. Part B gives basic data and references used in the preparation of the subsequent parts. Finally, parts C, D, and E contain the absorption coefficients in the ranges 1000°K to 24,000°K (Part C), 1 eV to 20 eV (Part D), and above 20 eV (Part E).

#6607005 8.153
Latimer, W. M.
OXIDATION STATES OF THE ELEMENTS AND THEIR POTENTIALS IN AQUEOUS SOLUTIONS SECOND EDITION

For each element the heats of formation, free energies of formation and entropies have been tabulated for all important compounds. Many of the free energies and basic enthalpy values have been estimated and a discussion has been given in the Appendix of the methods by which accurate estimations of entropy values have been made. Potential diagrams are given for all the elements which have more than one oxidation state. A discussion of the use of these diagrams in the interpretation of the chemistry of an element is given in Chapter 1. Several hundred new potential values and equilibrium constants have been added. The treatment of reaction rates and reaction mechanisms has been extended. The chemistry of uranium, neptunium, plutonium and americium has been greatly amplified and the actinide elements are treated in a new chapter. For the great majority of substances, the U. S. Bureau of Standards values for reaction heats have been employed.

#6902001 8.153
Leland, T. W.; Rowlinson, J. S.; Sather, C. A.
THE STATISTICAL THERMODYNAMICS OF MULTICOMPONENT SYSTEMS OF MOLECULES OF DIFFERENT SIZES

The central problem in the theory of mixtures is the calculation of the free energy of mixing of molecules of different sizes. An explicit calculation of this free energy is made for a mixture in which all intermolecular potentials are of the form $u(r) = \frac{A}{r^n}$, where $A$ is a distance characteristic of the interaction of two molecules of species $\alpha$, and where

$$\delta g = \frac{1}{2}(\gamma_{\alpha\beta} - \gamma_{\alpha\alpha} - \gamma_{\beta\beta})$$

when $\alpha \neq \beta$. This result follows from a solution of Percus-Yevick's integral equation for the pair distribution function of a mixture of hard spheres.

The form of the free energy provides a criterion by which existing theories can be judged, and it is shown that an approximation of the type originally suggested by van der Waals is superior to approximations based on the concept of random mixing. Molecules that differ only in size mix with a small and negative excess free energy. This recommended approximation is confirmed by comparison with experiment.
This text reviews the laws and uses of thermodynamics and includes tables and graphs of thermodynamic properties, constants, and applications. The tables include properties of normal fluids and gaseous solutions, entropy change in translation of an ideal gas to liquid, data for aqueous electrolyte solutions, Debye functions for the thermodynamic properties of solids, tabular summary of thermodynamic formulas, thermodynamic properties, electromagnetic work, symbols, and physical constants and numerical factors.

Lick, W. J.; Emmons, H. W.*8.153
THERMODYNAMIC PROPERTIES OF HELIUM TO 50,000°K Cambridge, Massachusetts: Harvard University Press 1962 122p

The thermodynamic properties of helium have been calculated at temperatures from 8,000°K up to 50,000°K and at pressures from 10⁻⁴ to 10⁹ atmospheres. In this range of temperatures and pressures, helium will consist mainly of a mixture of neutral atoms, singly and doubly ionized atoms, and electrons. Two Mollier enthalpy-entropy diagrams are included.

Liley, P. E.*8.253
COLLISION INTEGRALS FOR THE LENNARD-JONES 6-12 POTENTIAL TPRC Report 15, Thermophysical Properties Research Center, Purdue University, Lafayette, Indiana February 1963 15p 987

Values are presented for collision integrals occurring in expressions for the transport properties of non-polar gases assuming the Lennard-Jones 6-12 intermolecular potential function. Values of g11 and g12 are given for 0.100(0.001) 1.200(0.005) 2.00(0.01) 5.00(0.05) 10(1) 20 = 7/8 where 0 = a/\kappa. An estimate is made of the accuracy of the tabulation.

Liley, P. E.*8.253

Recent theoretical and experimental studies of transport properties of gases and gas mixtures are reviewed. Emphasis is given to data reported for high temperatures while no data for temperatures below the ice point are reviewed. Summary tables are given to enable easy access to references for the pressure and/or temperature dependence for a particular property and gas. Suggestions are appended for work urgently required and a bibliographic listing of 55 references given.

Linke, W. F.*8.253

Volume I, completed in 1957, includes material abstracted through 1956 and covers the elements and their compounds, alphabetically, from Argon (A) to Iridium (Ir.). The various types of solubility data are arranged in the following order: solubility in water; in aqueous acids; in aqueous bases; in aqueous salts having a cation in common with the solute; in aqueous salts having no common ion; in aqueous solutions of organic compounds; in anhydrous organic solvents; in high temperature mixtures (salt melts). The second volume of Solubilities completes the fourth revision of tables of Inorganic and Metal-Organic Compounds. It includes the solubilities of compounds from K through Z.

Lucks, C. F.; Deen, H. W.*8.153

Thermal conductivity, linear-thermal-expansion, specific-heat, density, and thermal-diffusivity values are reported over the temperature ranges shown for the following metals: 2024-T4 and 7075-75 aluminum from -250 to 800°F chromium from 75 to 3000°F, copper from 75 to 1900°F (density from 75 to 3000°F), Inconel and Inconel "X" from -250 to 1800°F, and AN-29 magnesium from -250 to 600°F, molten Xerox E from 75 to 3000°F, and diffusivity from 400 to 1600°F, "X" monel from -250 to 1800°F, SAE 1010 steel from -250 to 1650°F, stainless steels types 501, 316, and 347 from -250 to 1800°F.

McBride, B. J.; et al.*8.253
THERMODYNAMIC PROPERTIES TO 6000°K FOR 210 SUBSTANCES INVOLVING THE FIRST 18 ELEMENTS NASA SP-3001 Lewis Research Center, Cleveland, Ohio 1965 323p
Available from CFSTI

Consistent tables of thermodynamic properties at temperatures from 0 to 6000°K were compiled for 210 gaseous and condensed species involving the elements with atomic numbers of 1 to 18. The tables give the following thermodynamic functions for the standard state: heat capacity at constant pressure C_p, sensible enthalpy H^0-H_p, entropy S^0, sensible free energy (F^0-H_p), and so, the sum of sensible enthalpy and chemical energy at 0°K, as well as values of the enthalpy changes and the logarithms of the equilibrium constants. The latter two functions are given for the reactions of formation of the substances from a set of assigned reference elements. Data were taken from these elements in their atomic gaseous state (H^0 and log K). The functions for most of the gases were generated from molecular data, whereas the functions of most of the condensed species are

#6704034 *8.153
Lewis, C. N.; Randall, M.

This text reviews the laws and uses of thermodynamics and includes tables and graphs of thermodynamic properties, constants, and applications. The tables include properties of normal fluids and gaseous solutions, entropy change in translation of an ideal gas to liquid, data for aqueous electrolyte solutions, Debye functions for the thermodynamic properties of solids, tabular summary of thermodynamic formulas, thermodynamic properties, electromagnetic work, symbols, and physical constants and numerical factors.
A new equation of state for helium gas with six adjustable constants is presented. This relation is adequate for the representation of the P-V-T data and for the calculation of the entropy and enthalpy for a range of temperatures from 20 to 300 K, with pressures to 100 atm. A comparison of calculated volumes with original data indicates an average arithmetic deviation of 0.07 percent and maximum deviations of 0.5 percent. A comparison of calculated pressures with original data indicates about the same average and maximum deviations. A temperature-entropy chart and a compressibility factor chart have been prepared from values calculated by this equation of state. Values of density, enthalpy, and entropy, tabulated with temperature and pressure as independent arguments.

Selected P - ρ - T data for liquid and gaseous neon have been fitted to an analytical function. The selected data are the available experimental data, supplemented in regions of pressure and temperature where the experimental data are not available by values calculated from the theory of corresponding states and a generalized equation of state. Deviation plots are given which indicate the accuracy with which the analytical function represents the experimental data. The uncertainty of the tabulated properties is also estimated. The equation of state and the zero pressure specific heat were used to calculate tables of density, entropy, and enthalpy for selected temperatures and pressures. These properties are also illustrated in temperature-entropy diagrams.

Limited experimental pressure-density-temperature data for neon were compared with the pressure-density-temperature surfaces of neon by the law of corresponding states. Thus a basis is provided for the extrapolation of pressure-density-temperature data to lower temperatures and higher pressures.
provided concerning material processing, with special emphasis on specifications. Material suppliers are given. The Data Book has been organized for the most effective use of the data by the design engineer. Volume I contains light metal alloys (aluminum, titanium, etc.) data, Volume I-A contains nickel-base alloys, refractory metals and alloys, and nonferrous metals and alloys (cobalt, copper, etc.) data, Volume 2 contains ferrous alloys data, Volume 3 contains nuclear materials (fuels, neutron controls, beryllium, etc.) graphs, refractory ceramics, adhesives and plastics data. The section for each individual material contains descriptive basic data sheets with pertinent data relative to properties, characteristic-related information, together with curve sheets for graphic presentation of the parameters.

#6803003 *8.153
Meyer, C. A.; et al.
THERMODYNAMIC AND TRANSPORT PROPERTIES OF STEAM COMPRISING TABLES AND CHARTS FOR STEAM AND WATER
New York: American Society of Mechanical Engineers 1967
328p

The volume includes nine tables and thirteen charts on steam and water. The data were calculated from the equations and adopted by the International Formulation Committee and published in its 1967 "International Steam Tables for Engineering Use," in turn, were derived from the "International Skeleton Tables of 1963" adopted by sixteenth member nations of the Sixth International Committee on Properties of Steam. The following thermodynamic properties are listed: 1. saturation pressure and specific volume, entropy and enthalpy for saturated water as well as changes upon superheating from 72°F to 150°F; 2. saturation pressure and specific volume, entropy and enthalpy for steam and water. Specific internal energies from 3200 to 0.1 psia; 3. specific volume, enthalpy, and entropy of superheated steam and compressed water for temperatures from 1500°F to 32°F in 10-degree intervals, and pressures of 0.12 to 15,500 psia and also in the critical region (300°F to 650°F and 2800 to 3000 psia) at 2°F and 20 psia steps; 4. enthalpy as a function of pressure from 0.12 to 15,500 psia, and of the entropy in steps of 0.01 Btu/lbm-°F; 5. entropy values for same pressure range and for enthalpy in steps of 10 Btu/lbm-°F; 6. values of specific volume, enthalpy and entropy of superheated steam at pressures of 0.12 to 1500 psia; 7. specific heat at constant pressure for superheated steam; 8. thermal conductivity and Prandtl numbers of steam and water from 32°F to 1500°F and pressures from 1 to 7500 psia.

#6906029 *5.153-8.153
Miledé, J. T.; Clausen, W. V.; Goh, G. C.
FREE RADICALS AS HIGH ENERGY PROPULSANTS
General Electric Company, Flight Propulsion Laboratory, Cincinnati, Ohio
1966
150p
1062p

This compilation gives data concerning the use of free radicals as high energy propellants as well as discussions on rocket propulsion metastable states, and assumptions and methods of calculation. Tabulations and figures are included for the H, H₂, F, F₂, H₂, N, N₂, He, N₂, H, H₂, CH₄, CH₃, H₂, and I₂ systems. Also included in the thermodynamic properties given for each system are: speed of sound in a gaseous mixture in which chemical equilibrium is maintained at all times, speed of sound in a gaseous mixture in which chemical equilibrium is not maintained at all times. Appropriate data are included for the reactant mixture, molar enthalpy, equilibrium constants, mass of a particle, molecular weight, specific or total entropy, specific volume, and heat capacities of mixtures. This work represents an essential contribution to the study of rocket performance parameters and equilibrium composition in mole fractions is also listed. These data are listed for a broad range of temperatures and pressures for each system.

#6706013 *4.153-8.153
Miledé; J. T.; Welles, S. J.
BORON
BS-151, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California 1967
251p

A state-of-the-art survey and a data sheet compilation of boron and its compounds. The thorough literature search uncovered 394 references which were evaluated for information and data on the (1) reduction preparation methods, (2) zone-refining and melting techniques, (3) crystal structures (4) electrical, optical, and magnetic properties, (5) thermal and thermodynamic properties, (6) chemical and mechanical properties, (7) metallurgy and mechanical properties, (8) applications, and (9) availability and suppliers. Major emphasis was the characterization of boron's semiconductive and acoustic properties with respect to potential semiconductor and delay-line devices. Detailed information on the various methods of preparing pure boron and the resulting crystal structures plus properties are summarized in tables.

#6905016 *4.153-8.153
Miledé, J. T.; Welles, S. J.
CHEMICAL COMPOSITION AND ELECTRICAL RESISTIVITY OF ALUMINUM ALLOYS
BS-161, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California April 1969
27p

All commercially available aluminum alloys are identified by means of their standard designations. Available electrical resistivity data for these alloys are presented in tabular form for each alloy class. Electrical resistivity data for pure aluminum are reviewed.

#6910018 *8.353
Moore, R. T.; Harrison, R. T.; Douslin, D. R.
METHOD: BIBLIOGRAPHY OF THE THERMODYNAMIC AND TRANSPORT PROPERTIES ABOVE 300K
135p

Available from the Bureau of Mines

This bibliography of the properties of the methane above 300K is similar to and complements the one prepared by the U. S. Department of Commerce National Bureau of Standards (Technical Note 367) on the thermophysical properties of methane at temperatures below 300K.

#6902008 *8.153-3.153
Muettetters, F. L.
THE CHEMISTRY OF BORON AND ITS COMPOUNDS
New York: John Wiley & Sons, Inc. 1967
699p

This is a text on the properties of boron and boron compounds containing ten chapters, each written by experts in the different phases of boron chemistry. The chapters discuss the element boron, compounds of high boron content, borates, boric acid, boron hydrides and halides, boron-nitrogen, carbon,
phosphorus compounds, and sulfur and selenium compounds of boron. Each chapter contains a large listing of references as well as tables of the chemical properties of boron compounds.

#670630
*Muller, A. H., Jr.; Ando, K. J.; Coogan, H. H.
MOSSBAUER EFFECT DATA INDEX 1958-1965
New York: John Wiley & Sons, Inc.
1966
351p

The Mossbauer Effect Data Index is an organized index to experimental research results relating to the Mossbauer Effect published through 1965. The index groups experiments first by nuclear transition (daughter isotope and gamma-ray energy), then by source material or host, and then by absorber (or scatter) material or host. Each experiment is described and summarized on an 80-column IBM card. The data listed for each experiment are: Mossbauer isotope and gamma-ray transition energy, host material in which the source atoms are embedded, source temperature, absorber, absorber temperature, line width, dip of the spectrum (percentage change in counting rate from on-resonance to off-resonance), shape of the absorption spectrum, quadrupole splitting energy (for three transition lines), 1/2 lifet ime of the absorption spectrum and a code for the bibliography reference. A summary sheet is presented for each isotope presenting simplified decay scheme, gamma-ray and magnetic half-life and total internal conversion coefficient of the Mossbauer transition, natural isotopic abundance, and magnetic and quadrupole moments. There is an index to references by topic and author.

#6500320
*Mullins, J. C.; Kirk, B. S.; Ziegler, W. T.
CALCULATION OF THE VAPOR PRESSURE AND HEATS OF VAPORIZATION AND SUBLIMATION OF LIQUIDS AND SOLIDS, ESPECIALLY BELOW ONE ATOMESPER, V. CARBON MONOXIDE AND CARBON DIOXIDE
Technical Report No. 2, Georgia Institute of Technology, Engineering Experimental Station, Atlanta, Georgia
August 1963
81p

The vapor pressure and heats of vaporization and sublimation of carbon monoxide have been computed at one degree intervals from the normal boiling point (80.633°K) to 14°K. The vapor pressure and heats of sublimation of carbon dioxide have been computed at one degree intervals from the normal sublimation temperature (194.604°K) to 40°K. The agreement obtained between the computed and experimental vapor pressure data for carbon monoxide in the liquid and solid region down to the solid transition point (61.544°K) suggests that the computed values may well represent the experimental data within the accuracy of the temperature scales used by the various investigators. The computed values of the vapor pressures (Table XIX) are therefore recommended for use as smoothed values, especially below the solid transition temperature where the limited experimental data are scattered and discordant. For carbon dioxide the computed values of the vapor pressures agree well with the best experimental data which have been obtained in the thermodynamic scale. The experimental data measured on the International Temperature Scale appeared to deviate in a manner which is consistent with the available known deviations of the International Temperature Scale from the thermodynamic scale. The values of the vapor pressure of carbon dioxide computed at one degree intervals (Table XX) are therefore recommended as a consistent set of values on the thermodynamic temperature scale.

#6500321
*Mullins, J. C.; Ziegler, W. T.
THE SYSTEM HELIUM-ARGON FROM 65° TO 140°K UP TO PRESSURES OF 120 ATMOSPHERES, CORRELATION OF AVAILABLE PHASE EQUILIBRIUM DATA
Technical Report No. 3, Georgia Institute of Technology, Engineering Experiment Station, Atlanta, Georgia
January 10, 1965
64p

The available phase equilibrium data for the helium-argon system have been examined critically and a correlation of the gas and condensed phase compositions are presented from 65° to 140°K at pressures up to 120 atmospheres. No data are available for the solubility of helium in solid argon, but the location of the three-phase line suggests that the condensed solid phase is very nearly pure argon. The results of the correlations are presented in tabular form at 5°K intervals with the compositions of the equilibrium gas and liquid phases expressed in terms of mole fractions. In addition, (K = y/x) factors are given for both argon and helium as a function of temperature and pressure. The estimated maximum uncertainty of the gas phase composition in terms of the argon concentration is believed to vary as follows: ±4 percent from 65° to 110° to ±12 percent at ±10 percent at 140°K. The estimated uncertainty of the liquid phase composition expressed in terms of helium concentration is believed to be ±2 percent up to 110°K and ±5 percent from 110° to 140°K.

#6500388
*Mullins, J. C.; Ziegler, W. T.; Kirk, B. S.
THERMODYNAMIC PROPERTIES OF OXYGEN FROM 20° TO 100°K
Technical Report No. 2, Georgia Institute of Technology, Engineering Experiment Station, Atlanta, Georgia
March 1962
101p

The thermodynamic properties of oxygen have been calculated at two degree intervals from 30 to 100°K using published experimental data. The calculations cover the pressure range from approximately 1000 mm Hg at 100°K to approximately 100 mm Hg at 30°K. They include both the saturated and superheated regions. The properties calculated were the vapor pressure, heats of vaporization and sublimation, quality, volume, enthalpy and entropy. The enthalpy and entropy of solid oxygen at 8°K were taken to be zero. Calculations of vapor pressure and heat of sublimation were extended to 20°K. The tabulated data were used to construct a temperature-entropy chart for the range 30 to 100°K, showing lines of constant pressure, constant volume, constant enthalpy, and constant quality.

#6500387
*Mullins, J. C.; Ziegler, W. T.; Kirk, B. S.
THERMODYNAMIC PROPERTIES OF PARAHYDROGEN FROM 1° TO 22°K
Technical Report No. 1, Georgia Institute of Technology, Engineering Experiment Station, Atlanta, Georgia
November 1, 1961
68p

The thermodynamic properties of parahydrogen have been calculated at one degree intervals from 1 to 22°K using existing thermal and equation of state data. The properties calculated include the vapor pressure, heats of vaporization and sublimation, entropy and enthalpy of solid parahydrogen at 0°K were taken to be zero. The enthalpy and entropy of the vapor have been computed from the saturation pressure to 100 mm Hg. The tabulated results are shown in a temperature-entropy diagram with lines of constant pressure, constant volume, constant enthalpy and constant quality over the temperature range 4 to 173
22°K. In the selection of the experimental data used, heavy reliance has been placed on the work of Woolley, Scott and Brickwedde, and Pope and Arnold of the National Bureau of Standards. A more critical analysis of all existing data might result in slight changes in the computed results. These slight changes are unlikely to affect the use of the computed results for engineering calculations.

The water flow rate through the tank was measured by a constant-head tank and a turbine flowmeter. The tank was filled with a solution of known concentration and temperature, and the flow rate was measured by a calibrated turbine flowmeter. The temperature of the solution was measured using a calibrated thermometer. The concentration of the solution was determined using a calibrated spectrophotometer. The data was then used to calculate the electrical conductivity of the solution.

Available from the Superintendent of Documents, GPO

**#600016**
*$3.153-8.153$
Nelson, R. D., Jr.; Lide, D. R., Jr.; Maryott, A. A.
SELECTED VALUES OF ELECTRIC DIPOLE MOMENTS FOR MOLECULES IN THE GAS PHASE
NSRDS-NBS-10, National Bureau of Standards, Washington, D. C., September 1, 1967
49p

This table revises, brings up to date, and extends the coverage on numerical values of dipole moments which was included in NBS Circular 537, Tables of Dielectric Constants and Electric Dipole Moments of Substances in the Gaseous State, prepared by Maryott and Buckley in 1953. A recommended value with an estimate of accuracy is presented for more than five hundred organic and inorganic compounds. Comments are given on the definition of dipole moment and principal methods of dipole moment measurement, as well as an exposition of the criteria employed in selecting the tabulated data.

**#7066010**
*$4.153-8.153$
Neuberger, M.
DIAMOND DS-153, Electronic Properties Information Center, Hughes Aircraft Company, Culver City, California 1967
146p

Tables of electrical properties of diamond are presented as follows: conductivity, resistivity, dielectric constant, Hall coefficient mobility and thermoelectric power. Emission data are shown as well as a wide variety of irradiation effects. Energy data given are energy bands, energy gap, electronic levels, effective mass tables, vibronic and phonon energies. Optical properties included are absorption, reflection, refractive index, magnetoelectric properties, Debye temperature and thermal conductivity. A summary of crystal structure, phase transitions, and diamond types is included.

**#600016**
*$3.153-8.153$
Niles, F. E.
FORMATION OF IONS IN IONIZED AIR/NRL-R-1448, Ballistic Research Laboratories, Aberdeen Proving Grounds, Maryland November 1969
54p

Available from DDC

Schematic representations are given for both the formation of positive ions and the formation of negative ions in ionized air. The formation of positive ions can be divided into three regions. Region I contains the ionization of neutral constituents and two-body reactions in dry air. Region II contains reactions involving ionic species which are formed by three-body reactions. Region III contains reactions requiring water vapor and shows the formation of the hydrated hydronium ions. The formation of negative ions proceeds predominantly through $\text{O}_2^-$, which accepts one electron from neutral molecules to $\text{O}_2^-$, opens additional channels for the formation of $\text{NO}^-$, an ion which is not attacked by atomic oxygen. Rate constants are given, including the experimental values on which they are based, for the reactions in the schematic representations.

**#600002**
*$3.163$
Parker, V. R.
April 1, 1965
60p

Available from the Superintendent of Documents, GPO

The available specific heat, heat of dilution, and heat of solution data for aqueous solutions of uni-ions, with the exception of dihydrogen, have been critically reviewed and tables of selected "best" values at 25°C prepared. In addition, the neutralization data have been critically reviewed in order to obtain the thermodynamic properties. The electrolytes reviewed include hydroxides, halides and oxides, nitrites and nitrites, fluorides, and oxides, cyanides, cyanates, thiocyanates, and permanganates of hydrogen, ammonium, and the methyl ammonium derivatives, silver, and the alkali metals.

**#600169**
*$3.163$
Parsons, R.
HANDBOOK OF ELECTROCHEMICAL CONSTANTS London: Butterworths 1959
113p

This handbook has values for fundamental constants, properties of ions, salts and solvents, and numerical values of some functions used in the theory of electrolytes. Some of the properties covered include activity coefficients, thermodynamic properties of HCl and NaOH, thermal properties of electrolytes, equilibrium constants in solution, molar volumes of electrolytes and their temperature and pressure dependence; properties of molten salts, equilibrium properties of electrolytes, viscosity and diffusion in electrolytes; conductivity of electrolytes; kinetic properties of electrolytes; and surface tension, dielectric constant and refractivity of aqueous solutions.

**#6007016**
*$4.163-8.163$
Pomerance, H.
BIBLIOGRAPHY OF SECOND AND THIRD ORDER ELASTIC CONSTANTS ORNL-RNC-9, Research Materials Information Center, Oak Ridge National Laboratory, Oak Ridge, Tennessee December 1968
27p

Available from CFSTI

174
This bibliography of second and third order elastic constants of monocristalline materials was compiled from the files of the Research Materials Information Center and from Physical Abstracts. The literature since 1964 has been surveyed. The references are arranged first by the year of publication, next by the first author's initial, and last by a serial number. The tables do not show numerical values. The first table is the second order elastic constants, the second table is the third order elastic constants, and the third table is the third order constants; in each the substances are arranged alphabetically by chemical symbol. If the number of the reference is not complete, the letters c and s do not appear, but incomplete sets of derivatives are noted. The fourth column shows the format, whether the data are graphical, tabular, or by algebraic formula. The last column is the reference number.

#6706018
Pourhal, N. J. M.
ATLAS OF ELECTROCHEMICAL EQUILIBRIA IN AQUEOUS SOLUTIONS
644p
The Atlas presents data characterizing the equilibrium conditions of chemical reactions involving water, hydrogen, and of the elements, in the form of potential-pH diagrams. By means of these diagrams, it is possible to predict, on a thermodynamic basis, for a given element, the equilibrium states of all the possible reactions between this element, its ions, and its solid and gaseous compounds in the presence of water.

#6500419
Powell, R. L.; Blanpied, W. A.
THERMAL CONDUCTIVITY OF METALS AND ALLOYS AT LOW TEMPERATURES, A REVIEW OF THE LITERATURE
NBS Circular 556, National Bureau of Standards, Washington, D. C., September 1, 1954
68p
Available from the Superintendent of Documents, GPO

An extensive compilation is given of the measured values of thermal conductivity for metals and alloys from room temperature down to approximately 0.1°K. The most complete data set is plotted in 48 graphs. The tables and graphs for the metallic elements and alloys are essentially complete for literature reference from 1900 to 1954. For comparison, several graphs and tables are given for some representative dielectrics.

#6500494
Powell, R. L.; Roder, H. M.; Hall, W. J.
LOW-TEMPERATURE TRANSPORT PROPERTIES OF COPPER AND ITS DILUTE ALLOYS: PURE COPPER, ANNEALED AND COLD-DRAWN
Physical Review 115:314-323
July 15, 1959
Experimental results between 4° and 300°K are given for (1), the thermal conductivity, electrical resistivity, and thermoelectric force and power of two high-purity copper, one annealed and one cold-drawn 26%; and (2), the electrical resistivity of a series of seven samples of high-purity copper cold-drawn between 0% and 29% elongation. The total electronic thermal resistivities each consist of three terms: the intrinsic resistivity, the imperfection resistivity, and a deviation term. The temperature and one percent intrinsic resistivity and intrinsic electrical resistivity vary as $T^{1.7}$ and $T^{1.4}$, respectively, contrary to the predictions of the usual transport theory using Bloch approximations and assumptions. The resistivity of pure copper is 1.545 ohm cm at 0°C. The increase in imperfection electrical resistivity is approximately linear with increase in cold-drawn elongation. However, the added resistivity is not independent of temperature (Materials's rule), but about twice as great at the ice point as it was at 4°C. The change in thermoelectric power with concentration is positive at the lower temperatures, but negative above 38°C. The Lorenz number does not approach the Sommerfeld value at the lowest temperatures, but flattens out to a value considerably smaller. A discussion of each of the various effects is provided.

#6800018
Powell, R. W.; Ho, C. Y.; Liley, P. E.
THERMAL CONDUCTIVITY OF SELECTED MATERIALS
168p
Available from the Superintendent of Documents, GPO
This compilation consists of the critical evaluation and analysis of the available thermal conductivity data on eleven metals and nine metallic compounds, mainly from the GPO, Bureau of Standards, Bureau of Research and Development, and Bureau of Mines at Washington, D. C. and Oxford, England, from 1954. The data were selected mainly for their potential applicability as reference standards or because of their technical importance. The temperature range for which values are given often exceeds that for which these values are known with a high degree of certainty. The metals included are aluminum (solid and liquid state), copper, gold, iron (Armco and pure), manganin, mercury, platinum, platinum alloyed with 40 percent rhodium, silver, and tungsten. The nonmetallic solids treated include aluminum oxide, heryllium oxide, Corning code 7740 glass, diamond, magnesia oxide, Pyroceran brand code 9606 glass, quartz, thorium dioxide, and titanium dioxide. Data in the literature for the following substances in both liquid and gaseous state were examined and evaluated in this compilation: argon, carbon tetrachloride, diphenyl, helium, hydrogen, m-terphenyl, p-terphenyl, toluene, and water. Graphs, tables, and references are given for each substance.

#6500035
Fraunznitz, J. M.
SOLUBILITY OF SOLIDS IN DENSE GASES
48p
Available from the Superintendent of Documents, GPO
The thermodynamics of solid-dense gas equilibria is discussed, and two techniques are described for calculating the solution of a solid component in a gas at high pressure. The first one is based on the recent empirical equation of state of Redlich which, in turn, is derived from Pitzer's generalized tables of fluid-phase volumetric properties. The second one is based on the Flubrand-Scatchard theory of solutions. Both methods give good semi-quantitative results but cannot accurately predict solubilities from pure-component data alone.

#6500352
Predvoditelev, A. S.; et al.
CHARTS OF THERMODYNAMIC FUNCTIONS OF AIR FOR TEMPERATURES OF 100° TO 12,000°K AND PRESSURES OF 0.001 TO 1000 ATMOSPHERES
Glendig Ridge, New Jersey: Associated Technical Services, Inc. 1962
24p
Translated from Russian
This publication contains graphs of the thermodynamic functions of air plotted as a family of curves as a function of temperature and pressure, pressure being taken as the parameter. Computations of a normal shock and are not given, and information on this property must be obtained from the equation of state. In addition, this publication describes an approximate method for computing a normal shock and presents universal curves showing the pressure and enthalpy ratios for the shock as a function of the Mach number. At the end of the hook charts of "enthalpy vs. entropy" are given for an analysis of isentropic flow.

This paper presents correlations for three transport properties: viscosity, thermal conductivity and diffusivity for cryogenic liquids and their mixtures. A method for applying the correlations to the prediction of mixture properties is proposed. The basis for the correlations is provided by the theory of corresponding states. Attention is restricted to simple liquids along the liquid-vapor saturation curve from the triple point temperature to temperatures close to critical. The quantum fluids helium and hydrogen have not been included.

Rabinovich, V. A. (Editor)
Predvoditelev, A. S. (Translator)

THERMODYNAMICAL PROPERTIES OF GASES AND LIQUIDS No. 1
IPST Cat. No. 5554, Israel Program for Scientific Translations, Jerusalem, Israel
1970
24p
Available from CPSSTI, 69-55991

This collection is the first of a series of systematic publications initiated by the GSSSD - the Russian equivalent of the U.S. NSRDS - designed to supply reliable information on the thermophysical properties of substances and materials and aimed mainly at the practical application of the reference data. This collection includes papers dealing with the theoretical calculation and experimental determination of the thermophysical characteristics of substances and materials. It also contains some papers devoted to methods of calculation and the derivation of equations of state for pure substances and gas mixtures. Unlike existing publications, the collection presents the reference data in a form convenient for use; their accuracy is evaluated, and the methods used to prescribe within which the recommended values may vary. The reliability of the data obtained by theoretical calculation methods is checked against well-tested experimental values or confirmed by the accuracy of the calculation method.

Radebaugh, R.
THERMODYNAMICAL PROPERTIES OF He²-He* SOLUTIONS WITH APPLICATIONS TO THE He²-He* DILUTION REFRIGERATOR
December 29, 1967
15p
Available from the Superintendent of Documents, GPO

The thermodynamic properties of liquid He²-He* solutions between 0 and 1.5°K are calculated by using the weakly interacting Fermi-Dirac gas model for He in He*. Certain experimental data below about 0.4°K are used to evaluate some of the parameters in the model. The properties of both He in He* and the total solution are calculated for concentrations of He² up to 30 percent. All experimental data agree very well with the calculated results, although little data exist below 0.4°K. The calculated properties are used to analyze the behavior of the He²-He* dilution refrigerator in both the continuous and single-cycle processes. The maximum heat absorption below about 0.04°K is found to be 82 T² joules per mole of He². The effect of an imperfect heat exchanger and He* circulation on the refrigeration capacity is discussed. An analysis of two new types of single-cycle processes is also given.
This issue covers plutonium and its compounds and includes a critical evaluation of the data on thermodynamic properties, densities, crystallographic structures, compounds with non-metals, intermetallic phases, phase diagrams, equilibrium diagrams and diffusion rates in the condensed state. This assessment includes all the data published up to the end of 1964.

**#6510034**  
Reedy, T. L. (Editor)  
ICRPG WORKING GROUP ON THERMOCHEMISTRY: PROCEEDINGS OF THIRD MEETING MARCH 17-18, 1965, VOLUME I  
CPA Publication No. 82, Johns Hopkins Applied Physics Laboratory, Chemical Propulsion Information Agency, Silver Spring, Maryland  
July 1965  
172p  
Available from DDC

This volume contains the unclassified presentations given at the Third Meeting of the Working Group on Thermochemistry, and is concerned with data of interest for rocket motor performance calculations. The 22 papers include reviews of the status of thermochromal data for the heavy element compounds and ionized species. Recent experimental progress is reported in the areas of matrix spectroscopy, mass spectrometry, high temperature vapor pressure measurements of light element compounds.

**#609001**  
Reid, R. C.; Sherwood, T. K.  
THE PROPERTIES OF GASES AND LIQUIDS THEIR ESTIMATION AND CORRELATION, SECOND EDITION  
1966  
640p

This book presents a critical review of the various estimation procedures for a limited number of properties of gases and liquids. These include critical properties, P-V-T and thermodynamic properties, vapor pressures, latent heats, heats of formation, free energies of formation, heat capacities, surface tensions, viscosities, thermal conductivities, diffusion coefficients, and equilibrium properties of mixtures. Comparisons of experimental and estimated values are shown in the form of tables for indication of the degree of reliability of the methods discussed. The book is designed to be used as a text for molecular physics or a reference source for engineers. A set of Appendixes summarizes material referred to in the chapters and contains a large volume of tabulated data as well as references to the literature.

**#692061**  
Renon, N.; Eckert, C. A.; Prausnitz, J. M.  
MOLECULAR THERMODYNAMICS OF SIMPLE LIQUIDS, PURE COMPONENTS  
TIFC Fundamentals, 6:52-58  
February 1967

Thermodynamic properties of simple liquids are calculated from an analytical partition function based on a modification of Prigogine's cell theory and on a three-parameter theorem of corresponding states. The partition function gives an excellent representation of the configurational energy and entropy, volume, coefficient of expansion, and compressibility. These parameters are a molecular size, a pair-potential energy, and a term closely related to nonelastic intermolecular forces. This last parameter is a nearly linear function of Pitzer'sacentric factor. The main application of this statistical thermodynamic treatment follows from its straightforward extension to liquid mixtures.

**#690820**  
Rinshaw, S. J.; Ketchen, E. E.  
CESTIUM-137 DATA SHEETS  
ORNL-4186, Oak Ridge National Laboratory, Oak Ridge, Tennessee  
December 1967  
27p  
Available from CFSTI

Data on Cs-137 are tabulated. Properties of the fuel forms CsCl, CsSO4, and cesium borosilicate glass are given. Twenty-seven references are included.

**#690821**  
Rinshaw, S. J.; Ketchen, E. E.  
CURIUM DATA SHEETS  
ORNL-4187, Oak Ridge National Laboratory, Oak Ridge, Tennessee  
December 1967  
52p  
Available from CFSTI

Data on the fuel forms of Cm-244 are presented. Properties of Cm-244 metal, Cm0 2, Cm0 3, CmF 2, Cm0 2, and Cm0 3 cernet are tabulated. Forty-nine references are given.

**#690822**  
Rinshaw, S. J.; Ketchen, E. E.  
STRONTIUM-90 DATA SHEETS  
ORNL-4188, Oak Ridge National Laboratory, Oak Ridge, Tennessee  
December 1967  
45p  
Available from CFSTI

Data are given on composition, specific power, radiation, critical mass, container compatability, thermophysical properties, mechanical properties, chemical properties, biological tolerance, and shielding for strontium-90 metal, titinate (SrTiO3), oxide (SrO), fluoride (SrF2), and orthotitanate (Sr2TiO4). The data are indexed to 53 references which are also included.

**#6905025**  
Roberts, R. W.  
SUPERCONDUCTIVE MATERIALS AND SOME OF THEIR PROPERTIES  
NBS Technical Note 482, National Bureau of Standards, Washington, D. C.  
May 1969  
129p  
Available from the Superintendent of Documents, GPO

This is a noncritical compilation of data on superconductive materials that has been extracted from a portion of the literature published up to early 1968. The properties concerned are composition, critical temperature, critical magnetic field, crystallographic data, and lowest temperature tested for superconductivity. The compilation also includes a bibliography, general reference review articles and a special tabulation of high magnetic field superconductors. This compilation supersedes NBS Technical Note 408 having the same title.

**#6607006**  
Robinson, R. A.; Stokes, R. H.  
ELECTROLYTE SOLUTIONS, THE MEASUREMENT AND INTERPRETATION OF CONDUCTANCE, CHEMICAL POTENTIAL AND DIFFUSION IN SOLUTIONS OF SIMPLE ELECTROLYTES, SECOND EDITION (REVISED)  
London: Butterworths  
1965  
571p

This book is restricted chiefly to properties that are fundamental to electrolytes: conductance, chemical potential and diffusion. Conductance constitutes the main characteristic whereby electrolytes are distinguished from other
solutions. Of the thermodynamic properties, the Gibbs energy is the most useful for the treatment of equilibrium conditions and emphasis has been placed on quantities such as activities and ionization constants, which are related in a simple way to the Gibbs energy. Considerable space is given to the treatment of diffusion in electrolytic solutions. The text deals primarily with aqueous solutions though some recent data have been included and non-aqueous solutions whenever possible. Extensive appendices and tables are given, including functions and constants useful in computations, as well as compilations of accurate experimental data for non-ideal solutions. A reasonably full account is also given of the experimental methods where the data were obtained, in order to illustrate the accuracy and the limitations of modern technique.

#6500341 \*8.273
Roder, H. M.; Goodwin, R. D.
PROVISIONAL THERMODYNAMIC FUNCTIONS FOR PARA-
HYDROGEN
NBS Technical Note 130, National Bureau of
Standards, Boulder, Colorado
December 1961
139p
Available from CFSTI, Pb 161 631

#6510032 \*8.273
Roder, H. M.; Weber, L. A.; Goodwin, R. D.
THERMODYNAMIC AND RELATED PROPERTIES OF
PARAHYDROGEN FROM THE TRIPLE POINT TO 100°K AT
PRESSURES TO 340 ATMOSPHERES
NBS Monograph 94, National Bureau of Standards,
Boulder, Colorado
August 1965
109p
Available from the Superintendent of Documents, GPO

Experimental programs on parahydrogen at this
laboratory have provided pressure-density,
temperature relations and heat capacities at
temperatures from 15 to 100°K and at pressures from
2 to 350 atm. The two types of data have been correlated to yield a consistent set of functions. The properties tabulated for selected isobars and isochores are temperature, volume of pressure, the isotherm derivative (\( \frac{6p}{6T} \)), the isochore derivative (\( \frac{6p}{6T} \)), internal energy, enthalpy, entropy, the specific heats at constant volume and at constant pressure, and the velocity of sound. Also presented are the derived Joule-Thomson inversion curve and some comparisons with normal hydrogen near 100°K.

#6500318 \*8.173
Roder, H. M.; et al.
ORBITABARIC DENSITIES OF PARAHYDROGEN, DERIVED HEATS
OF VAPORIZATION, AND CRITICAL CONSTANTS
Cryogenics, 3:16-22
March 1963

Closely spaced experimental data are presented and used for defining a constant set of values for orbtobaric density, heat of vaporization, and critical constants of parahydrogen. Discussed are units and constants, experimental and derived data, and the various considerations affecting the solution of the critical temperature. Such

constants are to aid the calculation of thermodynamic functions in the single-phase region near the critical point.

#6500420 \*8.173
Rogers, W. M.; Powell, R. L.
TABLES OF TRANSPORT INTEGRALS
NBS Circular 595, National Bureau of Standards,
Boulder, Colorado
July 1958
46p
Available from the Superintendent of Documents, GPO

The transport integrals, \( J_n(x) \), defined by

\[
J_n(x) = \sum_{k=0}^{\infty} \frac{x^k}{k!}
\]

are utilized often in the development of theories describing the thermal conductivity, electrical conductivity, thermal electromotive force, specific heat, and other similar transport properties of solids. The tables include values to six significant figures of the integral for the index integral from \( x = 0 \) to \( x = 1 \) in the limit of integration \( x \) ranging from 0.1 by 0.1 intervals, to the limiting upper value, that ranges from 25 for \( n = 2 \) to 40 for \( n = 17 \). The three series utilized to represent the integral in different ranges are derived. The limiting values in the upper range and the asymptotic series expansions in the lower range are also given. Auxiliary tables include the values of the Riemann zeta numbers and Bernoulli numbers utilized in the calculations.

#6500390 \*8.273
Rossini, F. D.; et al.
SELECTED VALUES OF CHEMICAL THERMODYNAMIC
PROPERTIES
NBS Circular 500, National Bureau of Standards,
Washington, D. C.
February 1, 1952
126p
Available from the Superintendent of Documents, GPO

The tables of chemical thermodynamic properties are divided into two series. Series I gives the heat capacity at constant pressure, the heat of formation, the heat capacity, and the logarithm of the equilibrium constant of formation at all 25°C. The heat of formation at 0°K, molecular formula, description of substance, crystalline form, isomeric name, solvate, acid, base, if in solution, physical state, and a list of specific references are also given for each compound entry. Series II tabulates the heat, temperature and entropy of transition, fusion, and vaporization (including values of pressure) as well as the molecular formula, description, name of processes studied, initial and final states, and a list of specific references for each compound.

#6500379 \*8.373
Rossini, F. D.; et al.
SELECTED VALUES OF PROPERTIES OF HYDROCARBONS
NBS Circular 461, National Bureau of Standards,
Washington, D. C.
November 1947
496p

This book includes, in bound form, all the tables of properties of hydrocarbons (and certain closely related compounds) which have been issued as of May 31, 1947, by the American Petroleum Institute Research Project 44. Some explanatory remarks, and tables of fundamental constants, conversion factors, and molecular weights, are also included.
This report consists of a review of work done on phase equilibria of mixed systems of rare earth and other oxides. "Rare earth" here includes the elements yttrium and scandium as well as the lanthanide series; only trivalent oxides have been considered. The results are presented in the form of phase diagrams in temperature and percentage composition and as tables of the lattice constants of oxide crystals.

The theory of liquids and liquid mixtures is discussed and thermodynamic data for liquid mixtures and pure liquids are given. An account of the thermodynamic properties of pure liquids, a development of reliable methods for the calculation of the thermodynamic properties of a liquid in equilibrium with its vapor, the equilibrium properties of liquid mixtures, and an interpretation of the experimental material in terms of the intermolecular forces are presented. Accurate measurements of the properties of pure liquids including water and mixtures of such liquids as argon, nitrogen, oxygen, and the lower alkanes are also included. Information is included for mixtures of complex liquids, the critical state, high pressure mixtures, intermolecular forces and the statistical thermodynamics of fluids and mixtures. Indexes and references are given.

This book contains data on the physical, technical, mechanical, chemical, and refractory properties of refractories currently used and is widely used in technical development and in solving the problems of modern engineering. Chapters I-V of the reference book give information of a general character on refractory compounds, data on their crystal structure, specific gravity, thermochemical, thermal, electrical, and magnetic, optical, mechanical, chemical and refractory properties. In Chapters VI-VII, the author has been forced to provide some data of the resistance of refractory compounds to the action of different chemical reagents and molten media, and to oxidation. Chapter VIII gives concise tables of information on current and prospective fields of application of refractory compounds in different branches of industry. The data are organized according to the properties of more than 600 different materials such as borides, carbides, nitrides, silicides, etc. (oxides are excluded). In parts of the chapter, the following sequence in order of classes of compounds has been adopted: metal-like borides, carbides, nitrides, silicides, phosphides, sulphides, carbonitrides, and metallic compounds.

The reference book gives the most reliable data and indicates the literature sources in which duplicate values have been obtained by different investigators. Included are 1337 references.
resistance is largely governed by phonon interaction (anharmonicity of lattice vibrations), particularly at higher temperatures. The theory is well borne out by experimental data on dielectric crystals. By postulating a characteristic temperature dependence for each of the component thermal resistances corresponding to individual scattering effects, using the elements of the theory for each scattering mechanism, the total result is obtained in an empirical manner, in order to provide a rational basis for the experimental results.

Schmidt, E. (Editor)

PROPERTIES OF WATER AND STEAM IN SI-UNITS, KJ,BAR; ORIGIN: 1-1000 BAR; INCLUDING A MOLLIER H,S-DIAGRAM AND A T,S-DIAGRAM

New York: Springer-Verlag New York Inc., Title No. 1495
1969
205p

These tables of the properties of water and steam were calculated in their entirety by using a set of equations accepted by the members of the Sixth International Conference on the Properties of Steam under the title "The 1967 IFC Formulation for Industrial Use". Deviations from the figures compiled by Shaffer and Turner for smoothing only in small areas of isobaric specific heat capacity. For the most part, the tables on transport properties were also based on internationally accepted sets of equations. The quantity symbols and unit symbols used are those found in international standards. These Tables are mainly intended for use in industry. They are edited in concurrence with the major engineering institutions in a large number of countries and represent a valuable contribution towards international cooperation. These tables take the place of a new edition of the former Version B (published in 1963) of the "VDI-Wasserdampftafeln".

Schneider, S. J.

COMPILATION OF THE MELTING POINTS OF THE METAL OXIDES

NBS Monograph 68, National Bureau of Standards, Washington, D. C.
October 1963
31p

Available from the Superintendent of Documents, GPO

A compilation has been made of the melting points of 70 metal oxides published prior to January 1963. Both the original melting point and the equivalent value listed in the International Practical Temperature Scale of 1948 are presented. Included in the survey is information on pertinent experimental details such as the method of temperature measurement, purity, furnace type, and environmental conditions.

Scott, D. W.; McCullough, J. P.

CHEMICAL THERMODYNAMIC PROPERTIES OF HYDROCARBONS AND RELATED SUBSTANCES, PROPERTIES OF 100 LINEAR ALKANE THIOOLS, SULFIDES, AND SYMMETRICAL DISULFIDES IN THE IDEAL GAS STATE FROM 0° TO 1000°K

Bulletin 595, Bureau of Mines, Bartlesville, Oklahoma
1961
60p

Available from the Superintendent of Documents, GPO

Fifty-six tables are presented which record thermodynamic data. The properties covered are the free energy function, heat content function, heat capacity (enthalpy), entropy, heat capacity, heat of formation, free energy of formation and log of the equilibrium constant of formation for the above compounds at selected temperatures between 0° and 1000°K.

Shaffer, P. T. B.

THERMODYNAMIC FUNCTIONS FOR CARBON DIOXIDE IN THE RANGE 40 TO 1000°C AND 1 TO 1400 BARS

UCRL-7168, California University, Lawrence Radiation Laboratory, Livermore. California
December 1962
52p

Available from CFSTI

Extensive measurements on the pressure-volume-temperature relations of carbon dioxide were carried out sometime ago by Kennedy (1954). Since then, selected values of the thermodynamic function of carbon dioxide have been published by Price (1955) and by Majundar and Roy (1956). Even though these tables cover the data quite well, it still seemed desirable to make available the most extensive tables which could be calculated from the experimental data. The results of these calculations are illustrated. Tables are given for the temperature range 40 - 1000°C and the pressure range of 1 to 1400 bars which cover the properties: specific volume, entropy, enthalpy, Gibbs free energy, internal energy and Helmholtz free energy.

Shunk, F. A.; Dowdy, R.; Johnson, V. E.

CONSTITUTION OF ALLOYS ANNUAL REPORT NO. 1, JUNE 1967 TO MAY 31, 1968

IITRI-6602-1, IIT Research Institute, Technology Center, Chicago, Illinois
May 31, 1968
177p

A phased consolidation and updating of Hanssen's Constitution of Binary Alloys and its two supplements are being performed. In the first phase, reviews are under consideration for those binary alloy systems wherein at least one member of the pair is one of the elements Ac, Ag, Al, Am, Ar, As, or Au. These reviews cover the literature which was encompassed by various abstracting services through December 1966. Reviews for 121 systems are presented with this report.

Shunk, F. A.

CONSTITUTION OF BINARY ALLOYS, SECOND SUPPLEMENT

1969
720p

This book is the Second Supplement to the publication Constitution of Binary Alloys by Hansen in the Materials Science and Engineering Series. This supplement is concerned primarily with data which became available during the three-year period 1967-1969. It is intended to be comprehensive for this period. Approximately 8,700 system references were consulted in the preparation of the reviews for the 1,702 binary systems which form the text.
There are 219 figures. No data were previously available for 313 of these systems.


Each table summarizes the data up to 1960 for the association of one particular ligand with all the metallic ions which have been studied in conjunction with it, as far as the compilers located them. Some data published in 1961-1963 have also been included. Methods of measurement, composition, and temperature of the media to which the data refer, appropriate stability constants for the various complexes formed and references to the original papers are given for each ligand-metal pair. Acid dissociation constants of the ligands are recorded by including the hydrogen ion among the metal as one of the cations with which the ligands associate. Redox equilibria are represented by including the electron as a ligand, and hydrolysis of the metallic ion is described by reference to the hydroxyl ion as one of the ligands. The tables are divided into two sections, the first dealing with inorganic ligands and the second with organic ligands.

Simonds, H. R.; Church, J. M. A CONCISE GUIDE TO PLASTICS, SECOND EDITION New York: Reinhold Publishing Corporation 1963 404p

This second edition brings up to date the text of the first edition of A Concise Guide to Plastics and encompasses a much larger field due to the expansion of the plastics industry itself. An objective of this book was to answer the questions asked at the 1961 Plastics Exhibit. The subjects discussed include basic chemistry of plastics, manufacturing techniques, stereospecific catalysts, ablative, action, borderline materials, forms of plastics, resin manufacture, compounding, processing, applications, production and prices, selection factors, and the business of plastics. One chapter is devoted to plastic manufacturers' statements and contains information about the company's materials, trade names and sales for fifty-two producing firms. Over fifty pages of tables of the United States Plastics Trade Names are included, as well as numerous graphs and tables of data concerning plastic properties.


Available from the Superintendent of Documents, GPO

The three-phase region at and near the triple point of parahydrogen is presented on the graphical coordinates of temperature and entropy. Isobars from 10 mm Hg to 340 atmospheres, temperatures from 11°C to 239°C and specific volumes covering the range of from 10.5 cc/gm to 15,000 cc/gm are included. The energy base of enthalpy and entropy are consistent with previous data published by this laboratory.


Available from CFSTI

Commercially available gas, liquid and chemical sources of hydrogen for use with fuel-cell batteries are compared. Cryogenic storage is shown to be the most efficient on both weight and volume bases. Chemical generators are suitable for applications requiring small quantities of hydrogen gas at infrequent periods. Compressed-gas cylinders are convenient when small quantities of hydrogen are desired. A bibliography of selected publications during the past five years is included.


These annual volumes contain abstracts and bibliographies of papers published in the fields of thermodynamics and thermochemistry, arranged by country and laboratory. Each abstract contains a short description of the quantities measured and the method used. Topics of the abstracts include heats of combustion, reaction and formation, thermodynamic properties of pure substances and systems, vaporization studies, and other non-calorimetric thermodynamic properties. An author index and substance-property index are included. Each volume covers material from the year previous to the publication of the volume. Volume No. 2 was published in March 1959 and No. 4 in March 1961.


These annual volumes contain abstracts and bibliographies of papers published in the fields of thermodynamics and thermochemistry, arranged by country and laboratory. Each abstract contains a short description of the quantities measured and the method used. Topics of the abstracts include heats of combustion, reaction and formation, thermodynamic properties of pure substances and systems, vaporization studies, and other non-calorimetric thermodynamic properties. An author index and substance-property index are included. Each volume covers material from the year previous to the publication of the volume. Volume No. 6 was published in April 1963 and No. 7 in April 1964.

Somayajulu, G. R.; Zwaalski, B. J. ADDITIVITY OF THE POTENTIAL BARRIERS ABOUT DOUBLE BONDS IN HALOGENATED ETHYLENES Chemical Thermodynamic Properties Center, Texas A&M University, College Station, Texas September 1, 1965 9p

Potential barriers about double bonds have been found to be additive of the contributions from pairs of bonds separated by C=C bond in substituted ethylenes. On the basis of this work, the torsional frequency of ethylene has been found to be 225 cm⁻¹. New torsional assignments have been suggested for some molecules.
A generalized procedure for predicting the energies of the paraffinic hydrocarbons is deduced on the assumption of (i) interactions between pairs of bonds attached to a carbon atom, (ii) interactions between trios of bonds attached to a carbon atom, and (iii) interactions between pairs of bonds separated by a C-C bond. The derived equation justifies Tatevskii's empirical approach and is shown to be not very different from Allen's equation without the steric terms. Under defined conditions this equation reduces exactly to Allen's equation. Barriers to free rotation were calculated in deriving the new equation. This new equation is further modified by including corrections for rotational isomerism in place of trigonal interactions. The new relation is better founded in principle than Allen's and gives more refined steric terms due to higher order interactions.

In Part II, the enthalpies of formation of the substituted methanes have been calculated on the basis of Zahn's model. A method has been developed for the calculation of the dissociation energy of the R'-R" bond in a substituted ethane. Based on the dissociation energy of the R'-R" bond, the enthalpies of formation of the substituted methyl radicals and the substituted ethanes have been calculated. The enthalpies of formation of the substituted methyl radicals can also be calculated on the basis of Zahn's model. In Part III methods have been developed for the calculation of the magnitudes of the bond and pair bond contributions based on relationships between bond energy and bond dissociation energy. The pair bond contributions for the pair of bonds C-C in C\_2H\_4, C\_2H\_6, and C\_3H\_6 have been determined. Also determined were the enthalpies of formation of C\_2H\_4 and C\_5 radicals, C\_2H\_5 radicals, C\_4H\_9 radicals, C\_4H\_10 radicals, C\_5H\_11 radicals, C\_6H\_13 radicals, and C\_7H\_15 radicals.

In Part IV the enthalpies of formation of the C\_2H\_5 type radicals obtained in Part III have been used to obtain the enthalpies of formation of the C\_2H\_5 type radicals. Based on a relationship between bond energy and bond dissociation energy, the dissociation energies of the R'-R" bonds in substituted ethylenes have been calculated. Using the bond dissociation energies and the enthalpies of formation of the substituted methylene radicals, enthalpies of formation of a few substituted ethylenes have been calculated. The calculated values have been found to be in good agreement with the observed values.
which water is one of the components. Tables 1695 et seq. are devoted to binary systems consisting of condensed other than water. In two tables (p1646-1667) details are given of equilibrium conditions in solid phases for systems with aqueous and non-aqueous solvents. The formulae of the different solid phases in equilibrium with the saturated solution for each compound listed are given and in addition to the transition temperatures, the solubility of the substance being investigated is shown.

#6602003 *8,175
Stern, K. H.; Weise, E. L.
HIGH TEMPERATURE PROPERTIES AND DECOMPOSITION OF INORGANIC SALTS PART I SULFATES
176p
Available from the Superintendent of Documents, GPO

The literature dealing with the high-temperature behavior of inorganic sulfates has been critically reviewed. Free energy functions of reactants and products of the decomposition reactions were calculated and have been tabulated from 298°C up to as high a temperature as possible. Free energy functions, equilibrium constants of reactions, and partial pressure of reacting gases have also been tabulated. Auxiliary data on phase transitions, densities, and kinetics of chemical decomposition have also been included.

#6912009 *8,173
Stern, K. H.; Weise, E. L.
HIGH TEMPERATURE PROPERTIES AND DECOMPOSITION OF INORGANIC SALTS PART 2. CARBONATES
32p
Available from the Superintendent of Documents, GPO

The literature dealing with the high-temperature behavior of inorganic carbonates has been critically reviewed. Free energy functions of reactants and products of the decomposition reactions were calculated and have been tabulated from 298°C up to as high a temperature as possible. Free energy functions and equilibrium constants of reactions were tabulated. Auxiliary data on phase transitions, densities, and kinetics of thermal decomposition have also been included. The literature of the endothermic decomposition kinetics of solids, as it applies to carbonates, has been reviewed.

#6500475 *8,143
Stewart, R. B.; Johnson, V. J. (Editors)
COMPRENDIUM OF THE PROPERTIES OF MATERIALS AT LOW TEMPERATURE (PHASE II)
WADD Technical Report 60-56, Part IV, Wright Air Development Center, Wright-Patterson Air Force Base, Ohio December 1961
500p
Available from DDC, AD 272-769

Phase II of the Compendium includes data sheets on compressibility factor, velocity of sound and entropy of fluids, vapor-liquid equilibrium concentration of binary mixtures of fluids, and electrical resistivity and thermal conductivity integrals of metallic solids. Data sheets are included for each of these properties for the following materials: compressibility factor (helium, hydrogen, neon, argon, nitrogen, xenon, krypton, argon, neon, hydrogen, nitrogen, helium); entropy (a T-S diagram for neon); velocity of sound (helium, hydrogen, argon, nitrogen, krypton, xenon, neon, helium); liquid-vapor equilibrium concentrations (helium in hydrogen, nitrogen, argon, methane; hydrogen in nitrogen, carbon monoxide, argon, methane); electrical resistivity (5% of the pure metallic elements); and thermal conductivity integrals (44 pure metallic substances, 36 non-ferrous alloys, 9 ferrous alloys and 4 glasses and plastics). In general, the data sheets present the data primarily in graphical form and in a mathematical model include tables of selected values, references to the sources of the data and other references. Appropriate comments of interest to the user are also given.

#6500323 *8,173
Stewart, R. B.; Timmerhaus, K. D.
THERMODYNAMIC PROPERTIES OF CRYOGENIC FLUIDS
33p
Available from the Cryogenic Data Center, National Bureau of Standards, Boulder, Colorado

Cryogenic Engineering Laboratory, National Bureau of Standards has undertaken a re-evaluation of the pertinent experimental data to provide tabulations of thermodynamic properties for advanced cryogenic systems design. Tables and graphs of properties, as well as analytic functions for computer programs, are now available for several of the cryogenic fluids. These tabulations are intended to provide the engineer with comprehensive and consistent sets of data which are hopefully the best representation of the existing experimental data. These compilations have also served to define the limitations of the existing data and to define the needs for new measurements. This paper summarizes the experimental data for the several cryogenic fluids and outlines some of the procedures utilized in providing the various sets of thermodynamic property values.

#6500406 *8,273
Strobridge, T. R.
THERMODYNAMIC PROPERTIES OF NITROGEN FROM 64 TO 300°C BETWEEN 0.1 AND 200 ATMOSPHERES
85p
Available from CFSTI, PA 161-630

The internal energy, enthalpy, entropy, and specific volume of molecular nitrogen are derived and tabulated as functions of temperature and pressure. In addition to a mathematical model for the pressure-volume-temperature surface, accurate functions are given for the representation of the vapor pressure, density of saturated liquid, specific heat of saturated liquid, and the specific heat at zero pressure. Tabular values in British units over the same pressure and temperature range are available as Supplement A of this Technical Note.

#6500405 *8,273
Strobridge, T. R.
THERMODYNAMIC PROPERTIES OF NITROGEN FROM 114 TO 540°F BETWEEN 1.0 AND 3000 PSIA, SUPPLEMENT A (BRITISH UNITS)
85p
Available from the Superintendent of Documents, GPO

The internal energy, enthalpy, entropy, and specific volume of molecular nitrogen are derived and tabulated as functions of temperature and pressure. In addition to a mathematical model for the pressure-volume-temperature surface, accurate functions are given for the representation of the vapor pressure, density of saturated liquid, specific heat of saturated liquid, and the specific heat at zero pressure. Tabular values in metric
units over the same pressure and temperature range are also available in Technical Note 129.

#6510042-6510045  *$8.273
Stull, D. R.
JANAF INTERIM THERMOCHEMICAL TABLES VOLUMES 1, 2, 3 AND 4
Midland, Michigan: Dow Chemical Company
August 1968
1800 looseleaf pages

The JANAF (Joint Army-Navy-Air Force) Thermochemical Data tables contain data on the thermodynamic properties of materials of interest in rocket propellant and related field calculations. The properties tabulated are: specific heat capacities, Gibbs energy and enthalpy, volume changes, and free energy of formation as functions of temperature. References to reports from which the data were taken are given for each compound. The compounds are arranged alphabetically by chemical symbol. Volume I (6510045) covers A thru B; Volume II (6510044) covers C thru CINO; Volume III (6510044) covers C1 thru Hg4; and Volume IV (6510043) covers I thru Z and the electron gas. In each case the formulas themselves are indexed by arranging the elements making them up in alphabetical order of chemical symbols.

#6510030  *$8.223
Stull, D. R.
JANAF THERMOCHEMICAL TABLES
Midland, Michigan: Dow Chemical Company, Thermal Research Laboratory
August 1965
1000 looseleaf pages
Available from CFSTI, PB 168370

Thermodynamic properties of the light elements and their compounds are given for the solid, liquid and gaseous state. Some of the properties covered include heat capacity, standard entropy, heat of formation, free energy of formation, free energy function, heat content, log of equilibrium constant, heat of fusion, heat of sublimation and vaporization, melting point, boiling point. Some references are given.

#6910028  *$8.273
Stull, D. R.
JANAF THERMOCHEMICAL TABLES, FIRST, SECOND AND THIRD ADDENDA
Thermochromical Research Laboratory, Dow Chemical Company, Midland, Michigan
215 100p, 100p
Available from CFSTI, PB 168370-1, PB 168370-2, PB 168370-3

These three addenda to the JANAF Thermochromical Tables represent the additions and revisions made by the Dow Thermal Research Laboratory during the period June 1965 to June 1968. The addenda represent a changeover to the C¹ atomic weight scale and to the physical constants recommended by the National Academy of Science-National Research Council and the National Bureau of Standards. A new element, copper, and some of its compounds have been added to the collection.

#6810010  *$8.173
Stull, D. R.; Sinke, G. C. (Compilers)
THERMODYNAMIC PROPERTIES OF THE ELEMENTS
Washington, D. C.: American Chemical Society, Advances in Chemistry Series 18
1956
233p

Tabulated values of the heat capacity, heat capacity entropy and heat free energy function of the solid, liquid and gaseous states of the first 92 elements are given for the temperature range 298 ° to 3000°K. Auxiliary data include temperatures and heats of transition, melting and vaporization, and vapor pressures. Literature sources are listed. The compiled values have been analyzed and are supplemented by estimates when experimental data are lacking.

#6011010  *$8.173
Stull, D. R.; Westrum, E. F., Jr.; Sinke, G. C.
THE CHEMICAL THERMODYNAMICS OF ORGANIC COMPOUNDS
New York: John Wiley & Sons, Inc.
1969
$35p

A comprehensive and systematic treatment of organic chemical thermodynamics is provided from a search of the literature to January 1, 1966. Tabulations have been limited to the temperature range from 298.15° to 1006°K. The heat capacity, entropy, enthalpy and Gibbs energy of formation for 741 pure organic compounds from 298 to 1006°K have been tabulated for the ideal gas state. The entropy, enthalpy, and Gibbs energy of formation are presented as full as possible for approximately 4400 organic compounds in the ideal gaseous and condensed states at 298°K. Some regularities existing between hydrocarbons and other classes of organic compounds are pointed out. A review of relevant elements of thermodynamics and a discussion of calculational methods are included in Section I. Brief descriptions of calorimetric measurements are provided and the primary sources of information are tabulated in Sections II and III. Chapter IV introduces the methods used to estimate thermodynamic properties of compounds that have not yet been measured experimentally.

#6511005  *$8.172-2.772
TABLES OF THERMODYNAMIC DATA, SUPPLEMENT TO THERMODYNAMICS OF NUCLEAR MATERIALS
Vienna, Austria: International Atomic Energy Agency
STI/PUB/39
November 1964
$95p
Available from UN Headquarters

A compilation is presented of data on the thermodynamic properties of materials of interest for nuclear technology. The data were released in the Symposium on Thermodynamics of Nuclear Materials, which was held in May 1962. The properties covered include: heats of formation, free energies of formation, entropies of formation, and standard entropies at 298 ° and at other temperatures; enthalpy increments, free energy functions, entropy increments; enthalpies, free energies, entropies of transitions; heats, free energies, entropies of reaction; heat capacities; vapor pressures; thermodynamic properties of mixtures; melting points, densities; surface tensions; and lattice parameters.

#6511006  *$8.172-2.772
THERMODYNAMICS OF NUCLEAR MATERIALS, PROCEEDINGS OF THE SYMPOSIUM ON THERMODYNAMICS OF NUCLEAR MATERIALS HELD BY THE INTERNATIONAL ATOMIC ENERGY AGENCY IN VIENNA, 21-25 May 1962
Vienna, Austria:International Atomic Energy Agency, STI/PUB/58
September 1962
$92p
Available from IAEc

The International Atomic Energy Agency initiated a project to assist in disseminating and assessing data on important nuclear materials. Thus, the symposium was organized in an attempt to present and appraise the results of the work on thermodynamics of nuclear materials; to assess their reliability and further to discuss the application of the results and principles of thermodynamics to the solution of current problems in nuclear technology. Particular emphasis was placed on the thermodynamics of actinides and vaporization processes.
The data compiled refer only to binary systems of concentrated solutions which include systems having 20 and 90 per cent by weight, except those where there is only one measurement between 10 and 20 per cent. All types of substances are considered as components with the exception of metallic alloys. The data are classified by systems, and are arranged by volume as follows: Volume I, both components are organic compounds, excepting the hydroxyl derivatives; Volume II, both components are organic compounds of which at least one is a hydroxyl derivative; Volume III, at least one of the components is a metallic compound; Volume IV, all other systems. Heterogeneous equilibria properties considered are: density, specific volume, vapor pressure for the triplicate equilibrium, vapor pressure curves, boiling point, composition of azeotropes, composition of liquid and vapor components, in liquid phase and in gaseous phase, critical constants, liquid density (solid), rectilinear diameter, composition of the two liquid phases and functionally of the saturated vapor, critical solution point, freezing and melting curves, magnetic and transition points, and equilibrium of the condensed phases under high pressure. Properties of the gas, liquid and mixed crystal phases discussed are: densities, critical constants, refractive index, density, surface tension, viscosity, and heat. The data have been taken as given in the original papers. A bibliography and compound and author indexes are included at the end of the fourth volume.
Methods used by the Thermophysical Properties Research Center for handling collected bibliographical information on thermophysical properties are described. Details are given regarding the method of coding abstracts for preparation on IBM cards. Special consideration is given to the coding of substances and the system of substance classification and notation used for this purpose. Procedures are described which are used to store the punched card information on magnetic tape. Systems for retrieval of this stored information are delineated including a description of the organization of the retrieved bibliography and of the machine techniques involved.


A listing of the properties of materials is presented. The properties listed are density, melting point, heat of fusion, heat of vaporization, heat of sublimation, electrical resistivity, specific heat at constant pressure, thermal conductivity, thermal diffusivity, thermal linear expansion, radiative properties (absorptance, emittance, reflectance and transmittance), and vapor pressure as a function of temperature. An explanatory section is included with each volume. The titles of the volumes indicate the type of materials included in each volume.

Touloukian, Y. S. (Editor) THERMOPHYSICAL PROPERTIES RESEARCH LITERATURE RETRIEVAL GUIDE, BOOKS 1, 2 AND 3, SECOND EDITION New York: Plenum Press 1967 819p, 625p, 1315p

All research literature from 1920 to 1964 has been searched for the thermophysical properties of given substances. The specific properties considered are thermal conductivity, specific heat at constant pressure, viscosity, thermal radiative properties, diffusivity, electrical resistivity, thermal diffusivity and Prandtl number. Book 1 contains a listing of substance, the properties compiled for each substance and the classification number for each substance. The Search Parameter Index, Book 2, is arranged by property and by substance classification number. For each substance, one or more compilation serial numbers are listed together with the information on thermophysical, subject of the publication, language, temperature range considered and year of publication of the reference. Book 3 contains the Master Bibliography and an Author Index.

URANIUM-CARBON AND PLUTONIUM-CARBON SYSTEMS, A THERMOCHRONOLOGICAL ASSESSMENT, A REPORT FROM THE PANEL ON THERMOCHRONOLOGICAL PROPERTIES OF THE URANIUM-CARBON AND PLUTONIUM-CARBON SYSTEMS HELD IN VIENNA, 8-12 OCTOBER 1962 Vienna, Austria: International Atomic Energy Agency, STI/DOC/10/14 February 1963 48p Available from International Atomic Energy Agency Thermodynamic data reported by various laboratories on U-C and Pu-C systems are reported. Efforts were made to resolve various discrepancies. Sections are included on preparation, labelling, thermodynamic phase diagrams, heat capacities and thermal properties of the actinide carbides, heat of formation of the uranium carbides, vaporization studies, high-temperature equilibriums measurements, and the Pu - C system.


This article reviews the spin waves method of calculating the magnetization for a crystal composed of regularly spaced atoms at low temperatures, where magnetization differs only slightly from that at absolute zero. This article assembles in one place and in a unified fashion many of the results which are rather scattered in the literature, and introduces a simplified version of the quantum-mechanical theory. The approach is based on approximating the magnetic spin system by a system of harmonic oscillators. The treatment is based on the Heitler-London model for a magnetic solid, which is analogous to the Heitler-London model of chemical bonds. The magnetism is regarded as coming entirely from electronic spins regularly spaced in the crystal.


Thermodynamic data in condensed media obtained by shock wave techniques are listed. Both dynamic variables and volumetric quantities associated with the shock front are given. The information tabulated is derived from the shock compression of a sample of known density. The measurable quantities tabulated are pressure, volume or density, free surface and particle velocity behind the shock front, as well as shock velocity. The properties of the sample before shock compression are listed if available. The volumes are divided into the following sections: Section A-1, elements; Section A-2, inorganic compounds, including intermetallic compounds; Section B, hydrocarbons, organic compounds containing only C and H; Section C, other organic compounds; and Section D, carbides, alloys, solutions and mixtures. A materials index is included as well as a source index for each section.
The thermophysical properties of liquid nitrogen, oxygen, argon, and air are examined in this monograph. This book supplements previous studies by the authors on the thermophysical properties of these substances in the gaseous state. The book surveys and analyzes the best-known equations of state for liquids. An equation of state is derived for liquids on the basis of the principles of statistical physics, and is valid for many temperatures and densities. A method is developed for setting up this equation; it is used to describe analytically the thermal properties of the four most important cryogenic liquids. The experimental thermal data for liquid oxygen, argon, and air are extrapolated to pressures of 500 bar. As a result, equations of state valid for pressures of practical importance have been established, and the thermal properties of these substances have been calculated for parameters which have not been studied experimentally. The values obtained for the density were also used in tabulating the viscosity and conductivity of the four liquids with the aid of the limited experimental material on transfer coefficients. It was thus possible to obtain all the data on the thermal and thermomechanical properties of liquid air for the pressures most important for practical applications, extending to the saturation and freezing curves.

In recent years, a series of measurements of the thermal conductivity were completed for steam over a range of temperature and pressure. Argon and nitrogen were both used as a control check during the course of the measurements of the thermal conductivity of steam. These gases, particularly nitrogen, have been investigated extensively and additional measurements were made for nitrogen and argon using the steam. The new data is given together with a correlation of much of the available data for both gases, as a function of temperature and pressure.

This handbook of fused salt systems is based on 800 sources from the literature in Russian, German, English, Italian, and Japanese. The systems are arranged alphabetically according to the elements involved. Information discussed in these volumes include methods of distillation, thermal analysis, visual observations of first appearing crystals, X-ray and optics of quenched phases, chemical analysis of various solid and liquid phases, the nature of solids and the heat of solution. The first volume deals only with binary systems and simple salts with a common ion. Systems with double salts as one of the components are given as a part of the corresponding binary system. The second volume includes salts without a common ion, ternary systems with a common ion, quaternary, and pentameric systems. The compilation shows methods of choice and gives data for thermal analysis of visual observations. When numerical data is not available or incomplete for construction of phase diagrams, graphs are given for selected transfer properties in order to find up desired quantitative value (such as position of liquid-ice) if necessary. Conversions in solid phase are explained when given.
substances and organic molecules containing not more than two carbon atoms. In some instances such as metal-organic compounds, data are given for substances in which each organic radical contains one or two carbon atoms. No values are given in these tables for metal alloys or other solid solutions, fused salts, or for substances of undefined chemical composition. The physical state of each substance is indicated in the column headed "State" as crystalline solid (c), liquid (liq), glassy or amorphous (amorp), or gaseous (g). Solutions in water are listed as aqueous (aq). For non-aqueous systems the physical state is that normal for the indicated solvent at 298.15 K.

Weatherford, M. D., Jr.; Tyler, J. C.; Ku, P. M. PROPERTIES OF INORGANIC WORKING FLUIDS AND COOLANTS FOR SPACE APPLICATIONS PROPERTIES OF INORGANIC ENERGY CONVERSION AND HEAT-TRANSFER FLUIDS FOR SPACE APPLICATIONS WADC Technical Reports 59-588 and 61-06, Aeronautical Systems Division, Wright Patterson Air Force Base, Ohio December 1959, November 1961 550p, 600p

These reports (WADC TR 59-588 and its revision WADC TR 61-06) are intended to serve as a properties handbook for various inorganic fluids which may have potential value as coolants or working fluids for space applications. The fluids are presented as four distinct classes: liquid metals, liquid nonmetals, and liquids. The liquid metals include mercury, cesium, rubidium, potassium, NaK(75), sodium, lithium, bismuth, and lead. The liquid nonmetals included sulfur, phosphorus, lithium hydride, and aluminum bromide. The gases include krypton, argon, helium, hydrogen and aluminum chloride. Data are presented, where available, for temperatures ranging from less than 600°F to 4500°F for lead, and for pressures ranging from less than one atmosphere to greater than twenty atmospheres. The enumerated properties include density, viscosity, thermal conductivity, specific heat, latent heats of fusion and vaporization, entropy, vapor pressure, critical properties, diffusion coefficients, surface tension, thermal neutron absorption and scattering cross sections, equilibrium species composition, melting points, ionization potential, dielectric constant, electrical resistivity, magnetic susceptibility, corrosion characteristics, and materials compatibility. The characteristics of the various fluids are discussed, and the recommended values for the fluid properties are presented in either tabular or graphical form, or both, with detailed documentation as to basis and source. Background material, including thermodynamic-cycle, heat-transfer, compatibility, and working fluid considerations is discussed. A summary of current research activities in this field is presented.


Available from the Superintendent of Documents, GPO.

This Bureau of Mines bulletin was prepared to compress in readily available form the heat content, heat of formation, and free-energy-of-formation data for 65 common elements and their respective oxides, halides, carbides and nitrides. The necessary theoretical chemical thermodynamics is discussed in part 1 of this bulletin. It is assumed that the user will be at least acquainted with the fundamentals of chemical thermodynamics. Accordingly, all derivations of the various functions and their interrelations have been eliminated. This part contains the essential formulas necessary for applying fundamental data to process problems, with the accompanying examples included mainly to illustrate their applications. A survey of literature published through 1950 permitted assembly of all accepted measured and estimated thermodynamic values. These values are included in this bulletin. To satisfy individual needs, these data have been presented in 3 separate forms: (1) Tables of heat content, heat of formation, and free energy of formation at various temperatures; (2) Equations relating the variation of the thermodynamic functions with temperature; and (3) Graphical plots of the variation of free energy of formation with temperature.

Wiencke, R. W. Wehrwein, H. I. (Translator) EXPERIMENTAL AND THEORETICAL DETERMINATION OF THE THERMAL CONDUCTIVITY OF THE PLASMA IN A HIGH-CURRENT CARBON ARC TPRC Translation 9, Purdue University, Thermophysical Properties Research Center, Lafayette, Indiana September 1960 24p

The total thermal conductivity for the plasma of a high-current carbon arc is determined from the observation of the cooling trend in the arc column and from a sudden circuit break in the arc current as a function of the temperature within a range from 4000° to 10,000°K, after which the total thermal conductivity is compared with the theoretically calculated values. The agreement between experimental and theoretical absolute values is satisfactory, whereas the two curves (kappa (T)) show a displacement relative to each other, in such a way that the curve found
experimentally is shifted toward temperatures which are lower than those of the theoretical curve. It is presumed that the cause of this phenomenon is to be sought in the existence of relaxation processes during recombination of atoms into molecules.

This set of four volumes deals with the semiconductor, semimetal and optical properties of the III-V compounds. Contributions from ten to fifteen scientists are included in each volume. The books are designed for reference work as well as texts on the graduate level. Volume 1 reviews key properties of the III-V compounds, with special emphasis on band structure, magnetic field phenomena, and plasma effects. In volume 2, the emphasis is on physical properties, thermal phenomena, magnetic resonances, and photoelectric effects, as well as radiative recombination and stimulated emission. Volume 3 is concerned with optical properties, including lattice effects, intrinsic absorption, free carrier phenomena and photoelectric effects. Volume 4 includes thermodynamic properties, phase diagrams, diffusion, hardness, and phenomena in solid solutions as well as the effects of strong electric fields, hydrostatic pressure, nuclear irradiation and nonuniformity of impurity distribution on the electrical and other properties of III-V compounds.

This second edition of The Optical Properties of Organic Compounds describes more than twenty-five hundred substances and nearly two thousand are placed on one or both of the two diagrams which are enclosed with the book for the determination of composition. All organic compounds whose optical properties were described prior to October 1, 1952 are included.

This chapter discusses the value of the principles of physical chemistry in the development of synthetic rubbers and their application particularly in the American synthetic-rubber industry in the period between 1940 and 1954. It presents information about the structure and properties of rubbers and some possibilities of altering the structure to produce new rubbers with desired properties. Much attention is given to the Government-produced butadiene-styrene copolymer, the GR-S, Neoprene, and other rubbers which were produced on a large scale during that period. A large amount of data concerning the physicochemical properties of these rubbers is tabulated as well as a list of 407 references to work that has been done in this area.

Twenty-nine commercial varieties of synthetic rubber are classified as (1) chloroprene polymers, (2) butadiene polymers (including co-polymers), the organic polysulfides, (4) isobutene polymers, (5) plasticized vinyl chloride polymers, and (6) dimethylbutadiene polymers. Familiar examples of each of these types are, respectively (1) Neoprene, (2) the German Ruma rubbers, (3) Thokol, (4) Vistanex, (5) Koroscal, and (6) "methyl rubber." Since the last type is chiefly of historical interest, it is treated only briefly. For each of the first five types a brief discussion is presented concerning the raw materials used, the chemical reactions during manufacture, and the general conditions of polymerization or condensation. There is also some discussion of compounding and vulcanization, and of the properties and uses of the product. Published data on the different types of synthetic rubbers are presented, together with comparative tables of values of density, refractive index, band structure, and properties of interest. In Germany and Russia synthetic rubber is used to displace natural rubber in all its applications; in the United States, especially, where natural rubber is notably deficient are of present importance. Synthetic rubbers show outstanding properties in temperature resistance to the effects of liquids, especially petroleum products, and in resistance to deterioration by light, heat, ozone, and oxygen. These superiorities offset the higher cost of the synthetic materials, which is in most cases at least three or four times that of natural rubber. A bibliography of over 200 references covers the literature of the decade 1930-40.
This report is a compilation of data on thermal radiative properties. It also includes a brief discussion of the basic fundamentals of thermal radiation and of the methods of measuring these properties. Much of the information has previously been distributed in DMC memoranda; however, it is consolidated in this report for the benefit of those with a broad interest in radiant heat transfer. Thermal radiative data are included for the following materials: titanium and its alloys; stainless steels; iron-, nickel-, and cobalt-base superalloys; the refractory metals (chromium, columbium, molybdenum, tantalum, and tungsten) and their alloys; coated materials for elevated-temperature service; and ceramics. Forty-one references are included.

Working Group on Infrared Backgrounds
INFRARED TARGET AND BACKGROUND RADIOMETRIC MEASUREMENTS, CONCEPTS, UNITS, AND TECHNIQUES
IRIA Report 2389-64-4, Michigan University, Institute of Science and Technology, Ann Arbor, Michigan January 1962 45p
Available from DDC

This report discusses concepts, units, and techniques for making and interpreting measurements of radiation from targets and backgrounds. Graphs, figures, and references are included.

Yatsimirskii, K. B.; Vasil'ev, V. P.
INSTABILITY CONSTANTS OF COMPLEX COMPOUNDS
New York: Consultants Bureau 1960 214p
Translated from Russian

This book gives the instability constants of 1381 complex compounds prefaced with an introductory section of a general theoretical nature examining methods of calculating instability constants from experimental data. As a result, conditions on the stability of complexes, and the main factors determining the stability of complex compounds in aqueous solutions. This compilation includes original and abstract literature through 1954 and some work published in 1955-56.

Younglove, B. A.; Diller, D. E.
THE SPECIFIC HEAT AT CONSTANT VOLUME OF PARAHYDROGEN AT TEMPERATURES FROM 15 TO 90°K AND PRESSURES TO 340 ATMOSPHERES
Cryogenics, 21:1-5 December 1962

The computation of thermodynamic properties, relative to that of ideal gas, of compressed fluid by means of the equation of state data requires the data to be extremely accurate. The uncertainties that are introduced through the derivatives of the equation of state would be minimized by supplementing the calculations with the constant volume specific heat data of the compressed fluid. Specific heats for both ideal gas and compressed liquid states, together with PVT data will permit computations around closed paths to check the overall consistency of the data. For these applications the current program on hydrogen properties includes calorimetric specific heats of parahydrogen at constant volume. There are reported in this paper measurements of \( C_v \) of the compressed fluid from well below to above the critical temperature.

Younglove, B. A.; Diller, D. E.
THE SPECIFIC HEAT OF SATURATED LIQUID PARAHYDROGEN FROM 15 TO 32°K
Cryogenics, 2:1-5 September 1962

The specific heat of saturated liquid parahydrogen \( C_v \) has been measured as part of a program for determining the thermodynamic properties of parahydrogen at high densities and low temperatures. Thermodynamic properties of the compressed liquid have been computed using \( C_v \) as the baseline. The specific heat of argon has been measured for comparison with that of Flubacher, Leadbetter, and Morrison. Also the \( C_v \) of parahydrogen has been measured by others, which provides an additional comparison. The results obtained in this work serve to evaluate the accuracy that may be expected in future measurements of compressed fluids with this apparatus.

Ziegler, W. T.; Mullins, J. C.; Kirk, B. S.
CALCULATION OF THE VAPOR PRESSURE AND HEATS OF VAPORIZATION AND SUBLIMATION OF LIQUIDS AND SOLIDS, ESPECIALLY BELOW ONE ATMOSPHERE. 1. ETHYLENE
Technical Report No. 1, Project A-460, Engineering Experimental Station, Georgia Institute of Technology, Atlanta, Georgia June 2, 1962 47p

A number of thermodynamic relations are presented for computing, or assessing the probable accuracy of, the vapor pressure and heat of vaporization and sublimation of liquids and solids as a function of temperature. The relations are exact and take into account gas imperfection in the vapor phase. The solution of the equations, in general, requires iterative procedures which can be readily handled on an automatic digital computer. Applications of certain of the relations to the computation of the vapor pressure and heat of vaporization and sublimation of ethylene from its normal boiling point (109.40°K) to 20°K are presented. The computed results are given at 2 degree intervals from 168 to 100°K and 5 degree intervals from 100 to 20°K. The calculated results are thermodynamically consistent with the available experimental data and represent a thermodynamically acceptable extrapolation of these data to 20°K.

Ziegler, W. T.; Mullins, J. C.; Kirk, B. S.
CALCULATION OF THE VAPOR PRESSURE AND HEATS OF VAPORIZATION AND SUBLIMATION OF LIQUIDS AND SOLIDS, ESPECIALLY BELOW ONE ATMOSPHERE. 2. ARGON
Technical Report 2, Georgia Institute of Technology, Engineering Experimental Station, Atlanta, Georgia June 15, 1962 43p

The vapor pressure and heats of vaporization and sublimation of argon have been calculated from 88° to 20°K using thermodynamic relations. The calculations made use of published thermal data for saturated liquid and solid argon and second virial coefficients of liquid argon. Comparison of the computed vapor pressures with available experimental vapor pressure data showed that the experimental vapor pressure data were not thermodynamically consistent with the published thermal and second virial coefficient data. However, it was found that by decreasing the assigned value of the heat of vaporization at the normal boiling point by about one percent, the computed vapor pressures agreed quite well with the experimental measurements of Flubacher et al., which appear to be the most accurate data. The
The vapor pressure and heats of vaporization and sublimation of argon have been calculated at 1 degree intervals from the normal boiling point to the triple point and at 5 degree intervals from the triple point to 20.4°K by a thermodynamic method. Two sets of calculations were made. One calculation, given in Table VII, made use of a selected set of "best" experimental values for the necessary physical and thermal data. The other calculation, given in Table VIII, made use of the same input data except for slight adjustments in the heat of vaporization and temperature of the normal boiling point. The calculated vapor pressures have been compared with the available experimental data. It was found that the computed vapor pressures agreed quite well with the smoothed experimental data of Brickwedde and Scott down to about 70°K (or 70°K) while progressively poorer, still appears to be within the uncertainty of the experimental measurements. There appears to be little basis on which to choose between the computed values given in Tables VII and VIII. Uncertainties in the "best" values of thermal data used, as well as uncertainty in the temperature scale corrections, no doubt still exist. However, since the computed vapor pressures given in Table VIII agree slightly better with the extensive and careful measurements of Brickwedde and Scott, the vapor pressures and heats of sublimation given in Table VIII are recommended for use. A table of vapor pressures interpolated to one degree intervals is included for convenience. The thermodynamic functions for solid methane in the ideal gas state at one atmosphere have been computed at two degree intervals from 90° to 200°K, assuming a rigid rotator, harmonic oscillator model, using a summation procedure for the rotational partition function which takes into account nuclear spin. These functions have been used in the calculation of the vapor pressure and heats of vaporization and sublimation mentioned above, assuming that no interconversion of the nuclear spin species occurs (i.e., the room temperature equilibrium state is "frozen in"). The vapor pressure and heats of vaporization and sublimation have also been computed assuming the classical statistical mechanical form of the rotational partition function. The rotational contribution to the ideal gas thermodynamic functions of methane (E, H, entropy) and (C, S) for the individual nuclear spin species A, E, and F and the "frozen" high temperature equilibrium mixture (A: E: F = 3: 2: 9) have been computed down to 50°K.
Thermodynamic relations have been used to compute the vapor pressure and heats of vaporization and sublimation of ethane at one degree intervals from 184.52 K (the normal boiling point) to 20 K. These results are presented in Table VIII. These calculations made use of the "best" available experimental data for the normal boiling point and triple point temperatures, the heat of vaporization at the normal boiling point and heat of fusion at the triple point, heat capacities of the saturated condensed phases, and a representation of the second virial coefficient as a function of temperature based on the Kihara core model. The necessary thermodynamic properties of the ideal gas phases were computed from statistical mechanical relations using an internal rotation barrier of 2875 cal/gm mole. Comparison of the computed vapor pressures with the available experimental data strongly suggest that only the vapor pressure data of Loomis and Walters and the smoothed data of the American Petroleum Institute Research Project 44 (which are based at least in part on the data of Loomis and Walters) are reasonably consistent with the thermodynamic data used to compute the vapor pressures. By variation of the internal rotation barrier height and the theoretical representation of the second virial coefficient computed vapor pressures which agree somewhat better with the data of Loomis and Walters than those given in Table VIII were obtained. However, in view of the uncertainty in the temperature scale of Loomis and Walters and the thermal input data (including the second virial coefficient) it is believed that the vapor pressures and heats of vaporization and sublimation given in Table VIII represent these properties of ethane as satisfactorily as they are known at this time. No comparison of the computed heats of vaporization and sublimation with experimental values was possible, since no measured values of those quantities appear to have been made. The thermodynamic functions (H°, H°, S° and C° of ethane in the ideal gas state at one atmosphere pressure have been computed at 10 K intervals from 20 to 300 K using an internal rotation barrier of 2875 cal/gm mole. They are given in Table XI.

New and revised values of physical and thermodynamic properties of hydrocarbons and related sulfur and nitrogen compounds of primary interest to the petroleum industry are presented in tabular form. Some of the properties include density; vapor pressure; boiling point and freezing point; refractive index; transition temperature; specific dispersion; viscosity, surface tension, compressibility factor; critical temperature, pressure and volume; density; heat content function; free energy function; standard entropy and enthalpy; heat capacity; heat of reaction, heat of formation; heat of combustion; free energy of formation; log of equilibrium constant of formation; heat of vaporization, fusion, transition; entropy of vaporization, fusion, transition; and free energy of vaporization. For each table of numerical values, specific references by author are cited. A periodically revised cumulative General List of References is included.
New and revised values of physical and thermodynamic properties for classes of compounds of interest to the chemical industry which are hydrocarbons and nonhydrocarbons including halogen, oxygen, and some heterogeneous nitrogen compounds and also many simple inorganic compounds. Some of the properties include density; vapor pressure; boiling point and freezing point; temperature of transition; surface tension; viscosity; refractive index; cryoscopic constant; critical temperature, pressure and volume; density; and compressibility factor; heat content function; free energy function; entropy; enthalpy, heat capacity; log of equilibrium constant of formation; heat of formation; free energy of formation; heat capacity of vaporization, of fusion and transition; entropy of vaporization, fusion and transition; and heat of vaporization, fusion and transition. For each table of numerical values, specific references by author are cited. A periodically revised cumulative General List of References is included.
NBS TECHNICAL PUBLICATIONS

PERIODICALS

JOURNAL OF RESEARCH reports National Bureau of Standards research and development in physics, mathematics, chemistry, and engineering. Comprehensive scientific papers give complete details of the work, including laboratory data, experimental procedures, and theoretical and mathematical analyses. Illustrated with photographs, drawings, and charts.

Published in three sections, available separately:

- Physics and Chemistry
  Papers of interest primarily to scientists working in these fields. This section covers a broad range of physical and chemical research, with major emphasis on standards of physical measurement, fundamental constants, and properties of matter. Issued six times a year. Annual subscription: Domestic, $9.50; foreign, $11.75*.

- Mathematical Sciences
  Studies and compilations designed mainly for the mathematician and theoretical physicist. Topics in mathematical statistics, theory of experiment design, numerical analysis, theoretical physics and chemistry, logical design and programming of computers and computer systems. Short numerical tables. Issued quarterly. Annual subscription: Domestic, $5.00; foreign, $6.25*.

- Engineering and Instrumentation
  Reporting results of interest chiefly to the engineer and the applied scientist. This section includes many of the new developments in instrumentation resulting from the Bureau's work in physical measurement, data processing, and development of test methods. It will also cover some of the work in acoustics, applied mechanics, building research, and cryogenic engineering. Issued quarterly. Annual subscription: Domestic, $5.00; foreign, $6.25*.

TECHNICAL NEWS BULLETIN

The best single source of information concerning the Bureau's research, developmental, cooperative and publication activities, this monthly publication is designed for the industry-oriented individual whose daily work involves intimate contact with science and technology—for engineers, chemists, physicists, research managers, product-development managers, and company executives. Annual subscription: Domestic, $3.00; foreign, $4.00*.

* Difference in price is due to extra cost of foreign mailing.

Order NBS publications from: Superintendent of Documents
Government Printing Office
Washington, D.C. 20402