This teacher's guide is for the second year of the Portland Project, a three-year integrated secondary science curriculum sequence. The first of two parts in this volume, "Motion and Energy," begins with the study of motion, going from the quantitative description to a consideration of what causes motion and a discussion of Newton's laws. There follows the development of the laws of conservation of momentum and energy, including a discussion of energy in biological systems, and culminating with a discussion of the kinetic molecular theory. Part two, "Chemical Reactions," builds basic chemical concepts necessary for the analytical approach in the third year course. Topics considered in this part include the mole concept, equation writing, energetics associated with chemical reactions, the dynamic nature of particles and their interactions, and the application of energy and equilibrium to chemical systems.

The Harvard Physics Project textbook and the Chemical Educational Materials Study (CHEMS) textbook are used for most of the reading assignments in the second year course. A review of the development of the three-year program, its rationale and content, and a three-year course subject outline are included in this volume. Notes to the teacher, examples of data, and problem calculations are included.
MOTION AND ENERGY CHEMICAL REACTIONS

PARTS ONE AND TWO OF AN INTEGRATED SCIENCE SEQUENCE

TEACHER GUIDE 1973 EDITION

prepared by THE PORTLAND PROJECT COMMITTEE
under a grant from THE NATIONAL SCIENCE FOUNDATION
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Dedication

This volume is dedicated to the memory of Vernon Cheldelin under whose guidance and leadership integration of the sciences for Oregon secondary school youth was begun in 1963.
Introduction

The Portland Project was initiated in the fall of 1962 when two secondary school teachers, one with background in CBA chemistry, the other having responsibility for PSSC physics, began to note and discuss the redundancy in their respective courses. Why should students be subjected to this repetitious and fragmented representation of the physical sciences? they asked. A Steering Committee met to pursue the problem further and perhaps enlist the support of a funding organization to permit its exploration in depth. Under the able and devoted leadership of Vernon Cheldelin, Dean of the School of Science at Oregon State University (deceased), two proposals prepared for support by the National Science Foundation were funded in the summers of 1963 and 1964.

Thirty-five scientists and teachers devoted various quantities of time as writers, consultants, pilot teachers, and evaluators, with the aim of ascertaining the feasibility and efficiency of the integration of chemistry and physics. Concurrently and subsequently, other groups in other parts of the country have carried on studies that are approximately parallel to this one. Though the conceptual development and points of emphasis differ, the various groups are satisfied that integration of science courses is not only feasible but highly desirable.

Dr. Michael Fiasca of the Education and Science Staffs of Portland State University conducted an evaluation which revealed that subject matter achievement in chemistry and physics and critical thinking abilities are enhanced among students who studied the integrated courses over those who study the separate disciplines of

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Federation for Unified Science (FUSE) was recently organized to act as a clearinghouse of information on integrated science courses. Victor Showalter at Ohio University is the chairman of this committee.
chemistry and physics. It should be emphasized that though these differences were apparent, it could not be demonstrated that they were statistically significant.\(^2\) A concomitant result showed that enrollments in the two-year integrated courses were dramatically greater than in the separate courses.

A survey completed April 16, 1967 showed that there were forty-four schools in twenty states using the Portland Project integrated chemistry-physics course.\(^3\)

Mounting evidence in the literature from prominent persons working in science education strongly supported this mode of organization. Dr. Jerrold R. Zacharias, the prime instigator of the PSSC physics program, exemplified the changing attitude of scientists and educators:

The division of science at the secondary school level, into biology, chemistry and physics is both unreasonable and uneconomical.

Ideally, a three-year course that covered all three disciplines would be far more suitable than a sequence of courses which pretends to treat them as distinct. Today such a three-year course would be difficult to fit into the educational system, but much of this difficulty might be overcome at once if such a course existed, and it might well be that present tendencies in education would soon overcome the rest.

In any case, a greater coordination of the three subjects is possible even within the existing framework. It is understandable that the groups which developed the existing programs, each of which faced great problems of its own as it worked toward its goals, were reluctant to embark on the larger task of giving coherence to the sum of their efforts. With the programs now complete or approaching completion, it may be that the time has arrived for this necessary step.\(^4\)

\(^2\) Detailed results of this study may be obtained by writing to Dr. Fiasca at Portland State University.

\(^3\) Detailed enrollment figures and addresses of people who are using the Portland Project courses may also be obtained from Dr. Fiasca.

\(^4\) From page 52 of Innovation and Experiment in Education, a Progress Report of the panel on Educational Research and Development to the U.S. Commissioner of Education, the Director of the National Science Foundation, and the Special Assistant to the President for Science and Technology, March, 1964.
Stimulated by the apparent success of their original work towards this kind of integrated course, persons close to the Portland Project began to discuss extension of their work to include biology with chemistry and physics in a three-year sequence. A third proposal was prepared in 1966 and granted support by the National Science Foundation. Dr. Arthur Scott, member of the Chemistry Department at Reed College who has had deep interest in the Portland Project since its inception, graciously offered his talents, energy and time to carry on the project after Dean Cheldelin's death.

A writing conference was conducted on the Portland State University campus during the summer of 1967 to develop materials such as teacher and student guides. Eight local pilot schools committed approximately five hundred students and twelve pilot teachers for testing and evaluation. Dr. Donald Stotler, Supervisor of Science for the Portland School District, has had an active part in this and other phases of this project.

Twenty-six persons whose functions were writing, consulting, analysis, and editing met on the Portland State campus beginning June 14, 1967 to begin preliminary work on the integrated course. Their first task was to formulate an outline that displayed logical content development utilizing concepts out of biology, chemistry and physics. Particular attention was paid to matching students' abilities, interest and maturity level with the sophistication of concepts as nearly as this was possible to do. Then the committee perused material developed by the national curriculum groups --PSSC, Project Physics, CBA, CHEMS, BSCS and IPS -- in search of material to implement the outline they constructed previously. In the absence of appropriate materials, major and minor writing projects were initiated.

The writing committee continued its work in the summers of 1968 and 1969 with Dr. Karl Dittmer, Dean of the Division of Science, as director. Four major projects were tackled and completed: (1) extensive revisions were effected in the three-year outline, (2) the first- and second-year courses were revised based upon
student and teacher feedback, (3) the third-year course was developed incorporating Harvard Project Physics materials as a main vehicle, and (4) an evaluation program for the three-year course was developed.
Working Committee

Carl Bachhuber - Physics
Portland State University
Portland, Oregon

Ray Ellickson - Physics
University of Oregon
Eugene, Oregon

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Portland, Oregon

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St. Louis, Missouri

Edward Fuller - Chemistry
Beloit College
Beloit, Wisconsin

Howard Browning - Chemistry and Physics
Benson Polytechnic School
Portland, Oregon

Curtis Guthrie - Chemistry
Wilson High School
Portland, Oregon

David Cox - Chemistry and Physics
Rex Putnam High School
Milwaukie, Oregon

Jean Halling - Chemistry
Beaverton High School
Beaverton, Oregon

William Devery - Psychology
Reed College
Portland, Oregon

Glen Hampshire - Chemistry and Physics
Jefferson High School
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Lois Helton - Editor
Beaverton School District #48
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Karl Dittmer - Co-Director and Biochemistry
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Marshall Herron - Physics
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Arleigh Dodson - Chemistry
Lewis and Clark College
Portland, Oregon

George Ingebo - Director of Research
Portland School District
Portland, Oregon

Len Dooley - Biology and Chemistry
Sunset High School
Beaverton, Oregon
Donald Stotler - Supervisor of Science
Portland School District
Portland, Oregon

Harold Wik - Science Coordinator
Beaverton School District #48
Beaverton, Oregon

John H. Wirtz - General Science
Portland State University
Portland, Oregon
Pilot Schools

The following schools have served as pilot school for the pilot course during one or more of the past three academic years.

Adams High School
Portland, Oregon
Glen Hampshire
Lloyd Meskimen
Thomas Miles

Aloha High School
Aloha, Oregon
Mary Lou Combs
Elvis Delling
Nelson Doeleman
Ted Parker

Beaverton High School
Roger Berg
Jean Halling
Lois Helton
H. Dean Smith

Benson Polytechnic School
Howard Browning
W. B. Chase
Robert Franz
W. L. Hoffman

Central Catholic High School
Portland, Oregon
Jacob A. Mosbrucker
Peter Roerig

Cleveland High School
Portland, Oregon
John Brown
Edmund McCollough

Franklin High School
Portland, Oregon
John Neeley
Joseph Sklenicka

Grant High School
Portland, Oregon
Myra N. Rose

Jefferson High School
Portland, Oregon
Ronald Kawamoto
Leslie Morehead
Kenneth Starbuck

Parkrose High School
Parkrose, Oregon
Donald Pearson

Rex Putnam High School
Milwaukie, Oregon
Dennis Axness
David Cox
Jerry Fenton
Henry Kilmer
Jack McGoldrick
Roosevelt High School
Portland, Oregon
   Renee Bergman
   Kenneth Fuller
   Sue Storms

Sunset High School
Beaverton, Oregon
   Leonard M. Dooley

Wilson High School
Portland, Oregon
   Curtis D. Guthrie
   Norman Sipple

Woodland High School
Woodland, Washington
   Donald G. Fry
   George L. Stroud
ACKNOWLEDGEMENTS

Linda Anderson - Receptionist
Cecelia Benner - Secretary
Judy Bierman - Secretary
Doni Budlong - Secretary
Alice Eyman - Secretary
Mary Grant - Artist
Ethel Hays - Secretary
Lothar Graudins - Artist
Neomia Kendrix - Technician
Peggy Munyer - Typist
Janice Page - Laboratory Assistant
Robert Poon - Laboratory Assistant
Dianne Poppino - Laboratory Assistant
Jean Sims - Secretary
Linda Tham - Laboratory Assistant
Jill Timmons - Secretary
Kathryn Ann Torvik - Artist
Shirley Tsang - Laboratory Assistant
Barbara Wiegele - Secretary

STUDENT PARTICIPANTS

Patrick Moore - Jefferson High School (sow bug experiment)
The decision to try to develop a three-year integrated science course which would replace the traditional three courses in biology, chemistry and physics is based on several considerations. Among them are:

(1) a conviction that modern developments have made the division of science under these three headings obsolete;

(2) a recognition that the traditional courses overlap in many areas, resulting in a great deal of duplication and repetition as in the gas laws, atomic and nuclear structure, calorimetry and the kinetic molecular theory;

(3) a feeling that terminal students, who take no more than one year of science, deserve to get a taste of all of science rather than just one aspect, as they do in the conventional programs; and

(4) a desire to emphasize the unity in the approach to natural phenomena and the similarity in the methods, techniques and apparatus used by scientists in all fields.

A natural question arises as to what distinguishes this course from a general science course expanded to three years. The answer is that this course does not consist of a number of unrelated topics that might be taken up in any order; rather, it treats science as a structure that proceeds from observation to the development of general principles and then to the application of those principles to more involved problems. The emphasis in a general science course is on the results of science; the emphasis here is on the methods and reasoning by which scientists have arrived at these results.

The three-year course outline shows that a number of topics such as properties of matter, energy, heat, and certain biological concepts are discussed at the first-year level and again later in the course. This re-cycling is deliberate. It is intended to introduce students in a semi-quantitative way to some of the
significant generalizations of science and to show how these generalizations arose. These topics are treated again in the second and third years when greater facility with mathematics on the part of students makes it possible for them to understand and appreciate discussions of these topics in the succinct and precise language of mathematics.

An excessively formal and quantitative approach is avoided in the first year for several reasons. Students at this level do not extract essential meaning from such a presentation of information; furthermore, first encounters with new ideas should proceed from an intuitive, non-quantitative confrontation to one that is more quantitative. Teachers have spoken out against teaching and learning methods which substitute equations, formulas and other quantitative representations for first-hand experience, word descriptions, examples and illustrations. These criticisms are just as valid for students who are very capable and very interested in science as they are for other students. Moreover, the mathematical sophistication of students at this level is such that they are unable to follow most mathematical arguments as explanations for natural phenomena.

The typical science experiences of most secondary school students consists of one or two years devoted to general science and biology. Few study physics and chemistry. A significant advantage to the course of study described here is that students are given a chance to study physics and chemistry at a level of rigor that is consistent with their ability and their mathematical maturity. Students who terminate their study of science at the end of one year get a significant exposure to the structure of biology, chemistry and physics as they are presented in the latest curricular developments. Students who might not elect science beyond the first year because of lack of interest in biology may be attracted by the chemistry or physics portions of the course and elect to take an additional year or two of science. Students who are "turned on" by biology may wish to pursue further study of biochemical topics in Years II and III.
FIRST YEAR COURSE

After considering these problems and goals, the general course outline for the first year of the course was derived. It consists of four main parts:

1. Perception and Quantification
2. Heat, Energy and Order
3. Mice and Men
4. Environmental Balance?

The year begins with a study of the perceiver, moves on to the perceived, and ends with the interaction of the perceiver with the perceived. The first-year student starts out by gaining a better awareness of the nature of his perception and senses -- the faculties that make him aware of the world around him. With an increased understanding of these perceptual abilities, he can turn to the environment and then relate himself to it. He finds that his perception is limited and that he often needs to call on technological and conceptual extensions and that even these have their limitations.

The importance of organization and classification as parts of perception is emphasized. The physical properties of matter are introduced and studied as aids in organization and classification of chemicals. The identification of unknowns by study of their physical properties and use of organized data on punch cards is the culminating experiment of the Perception unit.

Apart from the great diversity exhibited in nature, which the scientist must organize in order to comprehend, certain unifying principles are essential for deeper understanding. The most powerful of these is the energy concept, which is explored in the "Heat, Energy and Order" unit in several of its ramifications -- physical, chemical and biological. The discussion begins by developing an experimentally important energy form, viz., heat. The macroscopic aspects of heat as embodied in calorimetry are related to the microscopic in terms of random molecular motion. This builds confidence in the idea of the atomic nature of matter, which
is essential to much of the unit. Various energy conversions form the vehicle for extending and generalizing the energy concept. Nuclear energy is developed in sufficient detail to underscore its environmental and social significance. Finally, the thermodynamic limitations and implications of energy conversion are explored, ending with a view of life as a supremely artful organizer in nature, a mechanism powered by energy which creates wondrous "local order", but always at the expense of influencing its environment.

The growth of a mouse colony carries the thread of the unit "Mice and Men." As the colony develops, students learn many things about the concept of population. The food and water consumed and products eliminated tie the mouse colony back to the unit "Heat, Energy and Order", and point ahead to the chapter on communities and to the unit "Environmental Balance?".

The cell concept is given prime position in this unit. It is used to enter topics on reproduction, embryology and maturation which are observed in the mice and other organisms. The mice selected for the original colony are such that an experiment in Mendelian genetics comes out of the observations students make as the colony develops. In most of the chapters man is an important organism and receives as much attention as the mouse, although the data are often secondhand.

A rather unpleasant fact that must be faced is that as our population increases, and human activities are directed towards increasing the standard of living for this population, strains are placed upon the environment. As students discover in "Mice and Men," the size of the community has a relation to both the quantity of the food, water and energy required and the quantity of waste products produced. To develop the concept of a closed system and point out the necessity for environmental management, an analogy between the earth and a spaceship is made. Students are then introduced by a multi-media approach to the nature of some of our common pollutants (with emphasis upon air, water, heat, noise and radiation)
as well as their effects. Following this students are encouraged to undertake a rather detailed study of a particular type or aspect of pollution. Emphasis here is placed upon student activity, which may take any number of forms. The culminating activity centers around discussion of these special studies together with the complex relations involved within the environment. It is hoped that out of these studies students will become aware of threats which exist to man's future on this planet.

THE SECOND YEAR COURSE

The second year of the course is considerably more quantitative in its approach than the first. This is the case because (1) the students are one more year along in their mathematical preparation, (2) the students who elect to take a second year of science are more likely to exert the effort to master more difficult topics, and (3) many of the quantitative aspects of physics and chemistry are basic to an understanding of molecular biology, which is an important part of the following year's work.

The second year consists of two parts:

(1) Motion and Energy

(2) Chemical Reactions

Year II begins with the study of motion, going from the quantitative description of motion to a consideration of what causes motion and a discussion of Newton's laws. There follows the development of the laws of conservation of momentum and energy, including a discussion of energy in biological systems. This section, which is primarily mechanics, culminates with a discussion of kinetic molecular theory.

Due to recent advances in both molecular biology and biochemistry, the descriptive approach to biology has gradually given way to one that is primarily analytical. It is now necessary, even on the high school level, for the serious biology student to have a more thorough understanding of those concepts normally
embodied in the "modern" high school physics and chemistry courses. The major objective of "Chemical Reactions" is to build some of those basic chemical concepts that are necessary for an analytical study of "The Chemistry Of Living Matter" and "Energy Capture and Growth."

The following subtopics of this section help in the realization of the major objective: Some of the topics discussed are the mole concept, equation writing, energetics associated with chemical reactions, the dynamic nature of particles and their interactions and the application of energy and equilibrium to chemical systems.

THE THIRD YEAR COURSE

Year III consists of four parts:

(1) Waves and Particles
(2) The Orbital Atom
(3) Chemistry of Living Matter
(4) Energy Capture and Growth

The underlying rationale of the third year is a study of energy and its importance to life. The first thrust is to build the orbital model of the atom using, as background, waves, electromagnetism and historical models of the atom. Once the orbital model is established as a representation of the localization and directionalization of electronic energy, structural models are built to show how biopolymers are spatially arranged and experiments are done to give evidence of energy relationships. With shape, size and energy relationships of molecules established, the DNA molecule is introduced. The culmination of this work comes in the final section when photosynthesis is considered. With this topic, much that has gone before is brought to a logical focus.

These topics are most appropriately placed in the third year of the integrated sequence after students have developed some facility with basic ideas
from chemistry and physics - e.g., quantitative knowledge about energy, mechanism of chemical reaction, equilibrium, rate of reaction, the photon and wave nature of light, electrical phenomena, and kinetic molecular theory. They should not now simply parrot biochemical processes such as photosynthesis and cell respiration but should understand the many chemical and physical principles which underlie these processes.

Time is allotted at the conclusion of Year III for individual investigation and studies.
Three-Year Course Outline

<table>
<thead>
<tr>
<th>TOPIC</th>
<th>REFERENCE</th>
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<td><strong>First Year</strong></td>
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<td>Part One: Perception and Quantification</td>
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<tr>
<td>I. Sensing and Perceiving</td>
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</tr>
<tr>
<td>II. Measurement, Distribution, Organization and Communication</td>
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<tr>
<td>Part Two: Heat, Energy and Order</td>
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</tr>
<tr>
<td>I. Heat</td>
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<tr>
<td>II. Temperature and Chaos</td>
<td>PP</td>
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<tr>
<td>III. Energy</td>
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<tr>
<td>IV. Nuclear Energy and Radioactivity</td>
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</tr>
<tr>
<td>V. Trends in Nature</td>
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<tr>
<td>Part Three: Mice and Men</td>
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<tr>
<td>I. Reproduction and Development</td>
<td>PP</td>
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<tr>
<td>II. Genetics</td>
<td>PP</td>
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<tr>
<td>III. Genetics and Change</td>
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<td>IV. Populations</td>
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<td>V. Ecology</td>
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</tr>
<tr>
<td>Part Four: Environmental Balance?</td>
<td>PP</td>
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</tbody>
</table>

* PP designation signifies materials produced by the Portland Project.
## TOPIC

### Second Year

#### Part One: Motion and Energy

<table>
<thead>
<tr>
<th>I.</th>
<th>Motion</th>
<th>HP*</th>
</tr>
</thead>
<tbody>
<tr>
<td>II.</td>
<td>Newton Explains</td>
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</tr>
<tr>
<td>III.</td>
<td>Multi-Dimensional Motion</td>
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<td>IV.</td>
<td>Conservation</td>
<td>HP</td>
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<tr>
<td>V.</td>
<td>Energy - Work</td>
<td>HP</td>
</tr>
<tr>
<td>VI.</td>
<td>Kinetic Theory of Gases</td>
<td>HP</td>
</tr>
</tbody>
</table>

#### Part Two: Chemical Reactions

| I.  | The Mole as a Counting Unit | PP   |
| II. | Combinations of Gases       | PP   |
| III.| A Useful Form of P=kDT      | PP   |
| IV. | Chemical Equations          | PP   |
| V.  | Electrical Nature of Matter | CHEMS+ |
| VI. | Basic Particles             | CHEMS |
| VII.| Energy Effects in Chemical Reactions | CHEMS |
| VIII.| Rates of Reactions          | CHEMS |
| IX. | Equilibrium                 | CHEMS |
| X.  | Solubility                  | CHEMS |
| XI. | Acid-Base                   | CHEMS |
| XII.| Oxidation-Reduction         | CHEMS |

* HP designates Harvard Project Physics material.
+ CHEMS designates material derived from the Chemical Educational Materials Study.
XIII. Stoichiometry

<table>
<thead>
<tr>
<th>TOPIC</th>
<th>REFERENCES</th>
</tr>
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<tbody>
<tr>
<td>Part One: Waves and Particles</td>
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<tr>
<td>I. Waves</td>
<td>HP</td>
</tr>
<tr>
<td>II. Light</td>
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<tr>
<td>III. Electricity and Magnetic Fields</td>
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<tr>
<td>IV. Faraday and the Electrical Age</td>
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<tr>
<td>V. Electromagnetic Radiation</td>
<td>HP</td>
</tr>
<tr>
<td>VI. The Chemical Basis of Atomic Theory</td>
<td>HP</td>
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<tr>
<td>VII. Electrons and Quanta</td>
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<tr>
<td>VIII. The Rutherford-Bohr Model of the Atom</td>
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</tr>
<tr>
<td>IX. Some Ideas from Modern Physical Theories</td>
<td>HP</td>
</tr>
<tr>
<td>Part Two: The Orbital Atom</td>
<td></td>
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<tr>
<td>I. Atoms in Three Dimensions</td>
<td>PP</td>
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<tr>
<td>II. Many-Electron Atoms</td>
<td>CHEMS</td>
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<tr>
<td>III. Ionization Energy and the Periodic Table</td>
<td>CHEMS</td>
</tr>
<tr>
<td>IV. Molecules in the Gas Phase</td>
<td>CHEMS</td>
</tr>
<tr>
<td>V. The Bonding in Solids and Liquids</td>
<td>CHEMS</td>
</tr>
<tr>
<td>Part Three: The Chemistry of Living Matter</td>
<td></td>
</tr>
<tr>
<td>I. Monomers and How They Are Built</td>
<td>PP</td>
</tr>
<tr>
<td>II. Polymers or Stringing Monomers Together</td>
<td>PP</td>
</tr>
<tr>
<td>III. Polymers in 3-D or The Shape of Things to Come</td>
<td>PP</td>
</tr>
<tr>
<td>IV. Where the Action Is--The Active Site</td>
<td>PP</td>
</tr>
</tbody>
</table>
Part Four: Energy Capture and Growth

I. Energy Capture
II. Energy Consumption - Metabolism
III. Metabolism and Genes

V. How Polymer. Make Polymers
VI. Genes, Proteins and Mutations
MOTION AND ENERGY
A Note to Teachers

This Guide is in no way meant to supplant other Teacher's Guides. It has been produced to help bridge gaps which result from the integration. It is imperative that the participating teachers become familiar with the material of all guides.

The experienced classroom teacher knows that time is always a vital factor. One objective of this integration is time. The teacher should search for that correct emphasis which permits him to cover each subject in depth and yet not be-labor it in such a way that future materials are not reached. A "development section" and a "teacher organization sheet" are included to help the teacher find that correct emphasis. The teacher should be particularly alert to the tendency to "bog down" early in the course at the expense of materials to come later.

Symbolism has been used in the teacher's guide. The following is a list of the symbols and their meanings.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
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</thead>
<tbody>
<tr>
<td>PSSC</td>
<td>Physical Science Study Committee</td>
</tr>
<tr>
<td>CHEMS</td>
<td>The Chemical Education Materials Study</td>
</tr>
<tr>
<td>PP</td>
<td>Portland Project</td>
</tr>
<tr>
<td>HP</td>
<td>Harvard Project Physics</td>
</tr>
<tr>
<td>TG</td>
<td>Teacher's Guide</td>
</tr>
<tr>
<td>CEx.</td>
<td>CHEMS Exercise</td>
</tr>
<tr>
<td>CExpt.</td>
<td>CHEMS Experiment</td>
</tr>
<tr>
<td>HPEX.</td>
<td>Harvard Project Exercise</td>
</tr>
<tr>
<td>HPExpt.</td>
<td>Harvard Project Experiment</td>
</tr>
<tr>
<td>R</td>
<td>Harvard Project Reader</td>
</tr>
</tbody>
</table>
Designed for each topic are many different activities, such as text material, demonstrations, transparencies, student activities, film loops, experiments, films, problems and reader articles. To utilize most effectively these various options a class may do one of three things: (1) select those options which are most practical or most effective for the class; (2) use nearly all options in carefully planned and prepared class meetings; or (3) let different students be responsible for different options. Each of these three methods may be used alternatively or any one might always be used or a compromise to fit the occasion and alternative available could be arranged. Each of the three methods has drawbacks, however: (1) if only certain options are chosen the time is more flexible but the opportunities for approach to a student on a given topic through various media is narrowed. (2) if all options are available and used, time for class discussion is limited and a strain is placed on the teacher for planning and pre-class set up. (3) if each option is used by teams of students, time must be made available for reports to the rest of the class, which is a problem with periods of less than 50 minutes.

We have therefore attempted to give an idea of priorities as we see them, based on activities which will carry the concept of the section. Additional activities are always helpful, and specific teacher background will help determine which activities may be chosen. As with any of the media, reading in the text may be considered as a resource, to be used when other activities don't convey information or as a supplement to the other activities. A possible priority for activities might be:

- laboratory experience
- demonstration
- student activity
- film loops
- film
- transparency
- text
- home desk lab problems
- reader articles
- lecture

A two week time cushion has been planned in addition to time allotted for the material in the course. This cushion may be used for post lab, pre lab, or seminar discussions and testing during the fourteen week period.
Errata For Harvard Project Physics Problems & Text

Unit I

Chapter I

SG 1.2d Inconsistency in distance symbols \( s_2 = d_2 \)
SG 1.2d Position result omitted \( d = 24 \text{m} \)
SG 1.2f Position result omitted \( d = 120 \text{ miles} \)
SG 1.5 \( 8.5 \text{ yr} = 2.70 \times 10^8 \text{ sec} \)
SG 1.17 \( v = 40 \text{ mi/hr} = .67 \text{ mi/min} \)
SG 1.18 The acceleration vs time graph may be constructed using either accelerations calculated from the given data or derived from the velocity vs time graph by the use of the slope of tangents at selected points.

Chapter II

SG 2.8 Part c is false because the distance travelled between images 3 and 4 is too great.
SG 2.9 Parts c and d are both true since \( \Delta d \) between successive intervals is reasonably constant.

Chapter III

SG 3.14a \( v_{\text{max}} = 7.8 \times 10^2 \text{ m/sec} \)
SG 3.16 If \( F \) is expressed in lbs., conversion factor is .22 lb/N
SG 3.18a \( F_{eb} + F_{\text{eb}} \)

Unit III

Chapter IX

SG 9.25 Corvette velocity = 35.2 m/sec
Corvette momentum = \( 5.15 \times 10^4 \text{ Kg-sec} \)

Chapter X

SG 10.8 Specific Heat of Lead = 0.03
SG 10.10 Students need to know the definition of power
& 10.11 and the watt, which are contained in paragraph 2, page 53. This is review from Year I.
SG 10.16 In TG expanded answer section "Energy required" is 270 Kcal.
Programmed instruction found useful with Year Two Materials


Harvard Physics, *Equations*, Parts 1, 2, 3.

Harvard Physics, *Kinetic-Molecular Theory of Gases*

Kass, Gerard A.,

Vol. 1 *The Atom*

Vol. 2 *The Chemical Bond*

Vol. 3 *Forces that Drive Chemical Reactions*

Vol. 4 *Kinematics and Equilibrium*

Vol. 5 *Acid-Base Chemistry*

Vol. 6 *Redox and Electrochemistry*


Suggestions for Laboratory Procedures

A laboratory is a place where scientists look at phenomena under controlled conditions. It is a place for serious work. Always prepare for an experiment by reading the directions in the manual before you come to the lab. Make a special effort to know all precautions.

Do only the experiments approved by your teacher. If you wish to do an extension (this is encouraged), check with your teacher. This general rule is for the safety of you and your fellow students. Laboratory safety is as much an attitude as a set of rules. The lab will become a safe place for investigation if the student continually uses common sense about his safety and the safety of others. If any accident does occur, report to your teacher. What seems a minor injury may have severe consequences.

You will be asked to write laboratory reports. Opinions concerning the content of these reports vary greatly. It follows that teacher judgment will determine the type of laboratory reports you are asked to write. The following ways to improve laboratory reports are to be taken as suggestions only.

1) Mistakes should not be erased. If there is room for the correction, the mistake should be crossed out without obliterating it and the correction made. If there is insufficient room, an extra piece of paper should be added.
(2) Spelling and punctuation are important. Sentence fragments should be avoided.

(3) The report should be carefully planned. It is best to know what type of observations should be sensed and, if possible, what regularities can be found. Planning will lead to the placement of items in a logical sequence in the report.

(4) The name of the experiment should be included.

(5) The date on which the experiment was done should be included.

(6) The names of all participants should be included and the name of the person who actually prepared the report should be designated.

(7) Some reports should include a simple statement or schematic diagram of the apparatus used in the investigation.

(8) Some reports will require a brief explanation of purpose and procedure. If these are given in the laboratory manual, they should not be included in the report. Copying items is "busy work."

(9) Nearly all experiments require taking measurements and subsequent collection of data. This must be carefully tabulated. If it is possible for you to make data tables before coming to the laboratory, you will have more time for observation, which is a major part of any laboratory experience.

(10) If computations are required to interpret results, they should be included in the report. However, if several computations of a similar nature are needed, they should be illustrated with a typical example. Mathematical equations, not arithmetical operations, should be shown.
If the investigation could be altered to get better results, a statement to this effect should be included.

If the investigation suggests extensions, these should be described.

Reading professional reports from magazines such as The Journal of Chemical Education and Scientific American should result in better reports.

Many times the most significant information about the experiment is to be found by graphing results. Whenever appropriate, graphs should be included in the report; they give a picture from which regularities can be sought. You will find the following suggestions very helpful.

(a) Always use a full sheet of graph paper.
(b) Position the ordinate and abscissa far enough from the edge of the paper to allow proper labeling.
(c) Assuming a relationship exists, the abscissa should represent the independent variable; the ordinate, the dependent variable. As an example: The distance of the gas pedal from the floorboard in an automobile would be the independent variable, plotted on the x axis, while the speed of the car would be the dependent variable, plotted on the y axis.
(d) Each axis must show units - e.g., cm/sec.
(e) Labeling an axis should run parallel to the axis.
(f) The scale of each axis should be chosen such that the functional plot covers most of the graph paper.

(g) The name of the graph, the name of the experiment and the date of the experiment should be suitably placed on the graph.

(h) When plotting data, draw a circle around each point or construct an error flag to indicate the uncertainty associated with the measurements.

(i) Draw the smoothest possible curve suggested by your data.
### Outline: MOTION AND ENERGY

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A. GRAPHS AND EQUATIONS

You may recall from your math classes the meaning of the words "variable," "functional relationship," "graphical analysis," etc. The purpose of the exercises you are about to perform is to help you understand what these concepts have to do with the world of experimental science, i.e., the world outside of mathematics in which you observe, measure and predict the behavior of real objects.

First let us review some of these ideas from the viewpoint of a scientist. In some experiments we measure more than one quantity simultaneously. If there is some sort of necessary connection between the varying quantities, that is, if there is some sort of law of nature or mechanism which relates changes in one quantity to changes in another, the quantities are called variables. For instance, in Year 1 you observed the motion of a pendulum. The position of a pendulum bob varies as time goes on. The distance of the bob measured along a ruler can be considered one variable (call it \( d \)), while the time
observed on a stop watch can be considered the other variable (call it $t$).

The fact that the two measured quantities have some well determined connection to one another is often stated by saying that one variable is a function of the other, i.e., that they are functionally related. (In the above example $d$ is a function of $t$.) There are several common ways of expressing this relationship: (1) a table can be made which shows values of one variable for many corresponding values of the other; (2) a graph can be drawn in which values of one variable are plotted on one axis and values of the other are plotted on the other axis; and (3) an equation can often be found which simultaneously contains the two variables. When a functional relationship has been expressed in one of these three forms, it is often possible to transcribe this information directly into the other forms. In these exercises we will concentrate on graphs and equations. In particular we shall show how to transform the information contained
in some simple graphs (such as might be obtained from experiments) into equation form.

In plotting graphs we consider one of the quantities to be the independent variable and plot it on the horizontal axis. The independent variable is usually thought of as the quantity whose value is imposed on the experiment from the outside. For instance, in measuring the distance traveled by a car, we find the length of time the car has been moving to be important. Time would be considered the independent variable in this example. The other quantity is thought of as representing the resultant behavior of the system and is called the dependent variable. In our example the distance traveled would be the dependent variable. The dependent variable is plotted on the vertical axis. A graph of the car's motion could appear as shown below (assuming it were traveling with a constant speed):

For the sake of simplicity it is recommended that students use only positive values for $x$ and that they use a minimum of 4 or 5 points for each graph.

When a functional relation is expressed in equation form you all know how to obtain a
corresponding graph. For practice you should try plotting the following equations: \( y = 2x \); \( y = 6/x; \) \( y = x^2 \); and \( y = x^3 \). Consider \( y \) to be the dependent variable and \( x \) to be the independent variable. Note the characteristic shapes of these graphs. They are of the type you shall be most concerned with in this exercise.

The reverse process, i.e., obtaining an equation when you already have the graph, may not be familiar to you. However, it is extremely important in experimental science. Our main task now is to learn how to do this in some important simple cases.

B. FROM DATA TO AN EQUATION

When experimental data is plotted directly on a graph, it may be possible to write the equation for the resulting curve. In the cases we shall be studying the process is two-fold. The first step is to discover the proportional relationship, if any, that exists between the variables. The second step is to determine the value of a proportionality constant which enables us to transform the statement of proportion into a bona fide equation.
The individual points from which the curves were drawn are not shown. The dots are covered over by the curves. In general when particular points are indicated on a graph they are meant to signify experimental points. In plotting an equation, however, it is poor practice to emphasize particular points as they are arbitrary and have no more significance than any other part of the curve.
B.1 PROPORTIONAL RELATIONSHIPS

The appearance of some typical curves suggests certain proportional relationships between variables. Several common general cases are illustrated below.

In the first graph $y$ is directly proportional to $x$. Symbolically this is written $y \propto x$. The plot is a straight line. (Therefore this sort of relationship is often called a linear relationship.)

The second graph illustrates a case where $y$ is in inverse proportion to some power of $x$. The case you are most familiar with is $y \propto \frac{1}{x}$, but proportionality to $1/x^2$, $1/x^{1/2}$, etc. would yield curves of much the same shape. Notice how the curve bows in towards the origin and approaches the axes at the extreme ends.

The third curve is obtained in the case
where $y$ is proportional to a power of $x$. This may be written $y = x^n$, where the exponent $n$ may have values such as 2, 3, etc., depending on the particular situation. The curve approaches the origin for small values of $x$, but rises very steeply for large values.

B.2 THE PROPORTIONALITY CONSTANT

You probably already know that two quantities which are proportional to one another may be linked together to form a correct equation by multiplying one side of the proportion by a constant of proportionality. If, for instance, $y = 1/x^2$, the corresponding equation could be written $y = k/x^2$, where $k$ represents the constant of proportionality. When $k$ has different values the graph would have the same shape, but it would pass through somewhat different points. When $k$ is evaluated and expressed in proper units, it has special

*You may have realized that the linear relationship illustrated in the first graph is actually a special case of the power relationship where $n = 1$. However, this is such an important case, and yields such a distinctive curve, that it is listed as a case all by itself.
physical meaning in many experiments. In the earlier graphs you plotted as an exercise, the value of k for the linear relationship was 2, for the inverse relationship was 6, and for the power relationships was 1. Try plotting those equations with other values of k, say 1, 3 and ½ respectively, in order to see what bearing changing the constant has on the curves.

C. A SIMPLE LINEAR CASE: DENSITY OF METALS

The physical interpretation of a constant of proportionality can be easily illustrated in the case of a laboratory exercise with which you are already quite familiar. Suppose you had a number of pieces of metal, say of gold, silver and aluminum, of various shapes and sizes. If you measured the mass and volume of each piece and plotted these quantities versus each other on graph paper, you would probably obtain straight lines similar to those shown on the next page. The basic meaning of the graphs is the same in all three cases, viz., the greater the volume of material, the greater the mass of material, in direct proportion. This is
of course exactly what we expect, and it is just as likely for one substance as another. But, although the graphs are all straight lines, they differ in one important respect. The lines slant at different angles to the axis. They have different slopes. The mass of gold increases at a faster rate when the volume of material is increased than does the mass of aluminum or silver. The constant of proportionality expresses this difference for us; its value is, in fact, the slope of the graph.

The slope tells us the ratio of the change in the variable plotted on the vertical axis and the corresponding change in the variable plotted along the horizontal axis. The slope
is sometimes referred to as rise/run, or designated as \( \frac{\Delta y}{\Delta x} \) (where \( y \) refers to the vertical axis and \( x \) to the horizontal). Note the calculation for gold.

\[
k(\text{gold}) = \text{Slope} = \frac{\Delta y}{\Delta x} = \frac{20 \times 10^3 \text{kg}}{1 \text{m}^3} = 20 \times 10^3 \frac{\text{kg}}{\text{m}^3}
\]

Check yourself in this skill by finding the slope of the lines in the silver and aluminum graphs and then comparing your answers to those given below.

\[
k(\text{silver}) = 10 \times 10^3 \frac{\text{kg}}{\text{m}^3}
\]
\[
k(\text{aluminum}) = 3 \times 10^3 \frac{\text{kg}}{\text{m}^3}
\]

For the examples we have been using in this discussion, the constant of proportionality is given a special name—density. As you probably remember from Year 1, density is a characteristic property of matter that tells
This experiment can be duplicated with commonly available equipment, either for demonstration purposes or for student laboratory work. In addition to the information in the student text, the following should be helpful if you wish to perform the experiment:

Source of power can be a LabVolt 0-45 V. power supply utilizing the DC output: heater--double element 5 amp immersion type wired in series (IFS equipment); Triplett multimeter (1000 ma scale). Connect the items in series.

Water is contained in styrofoam cups during heating. (Two cups will be necessary, as the heaters should be submerged during operation. The second cup of water can be used for backup data.) It is wise to fit each cup

us how the mass of any particular sample of matter is related to its volume.

In summary, the step by step analysis of our graphical data in these examples leads us from a statement of proportionality to a mathematical equation and thence to an evaluation of density. For gold:

\[ m \propto v \]

\[ m = kv \]

where \( k \), the slope of the graph, is the density:

\[ \frac{20 \times 10^3 \text{kg}}{m^3} \]

D. ANOTHER EXAMPLE: ELECTRICAL HEATING

D.1 GETTING THE DATA

In the Year 1 you performed experiments in which the temperature of water was raised using electrical immersion heaters. Electrical energy was put into a water bath and the consequent rise in temperature was observed. You might recall that some of the ideas you were concerned with were conservation of heat, the meaning of a calorie and the conversion of energy. We will now go back and look at the data from these experiments and further analyze them in terms of graphs and equations.

In the Year 1 experiments, styrofoam tanks were used to hold the water and the immersion
heaters were operated from a direct current (D.C.) power supply with provisions for varying the voltage and current. We varied the mass of the water, the length of the heating time, and the current flowing through the immersion heater, while measuring the temperature change in the water. Time was measured with an accuracy of ± 5 sec. using the sweep second hand on a wrist watch. The mass of the water was found indirectly from a volume measurement. The temperature change was measured with a Centigrade thermometer, the accuracy being ± 0.2°C. The current flowing through the heating element of the immersion heater was measured with an ammeter to within 0.01 ampere. Perhaps you would like to repeat this same experiment with equipment readily available in your laboratory. Some typical results are given below in tabular form.

Data points show greater deviation when short time intervals and small masses are used. As longer heating times are used, the heating curves begin to show a plateau. We found this to be true for approximately 8-10 minutes of heating time using the smaller masses of water. The plateau is apparently due to an equilibrium heat exchange being reached between the coil and the water and the water and the environment. You may wish to point this out to students, since it will be observed to a greater or lesser degree on all the graphs. We feel that this lends a realistic factor to the analysis.

with a cardboard lid to reduce evaporation and heat losses. Holes may be cut for the heater and thermometer. Be sure to stir regularly and remove the bubbles from the immersion heaters. It would lend additional accuracy to the experiment if the temperature intervals extended above and below room temperature by the same amount. This is very time consuming, and in actuality the test runs were made in consecutive fashion, without changing the water baths.

perhaps you would like to repeat this same experiment with equipment readily available in your laboratory. Some typical results are given below in tabular form.
Temperature Change (°C) upon Heating Using 0.40 Amp Current

Table D.1

<table>
<thead>
<tr>
<th>Mass (kg)</th>
<th>Time (sec)</th>
<th>0.300</th>
<th>0.250</th>
<th>0.200</th>
<th>0.150</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>0.4°</td>
<td>0.6°</td>
<td>0.7°</td>
<td>1.1°</td>
<td></td>
</tr>
<tr>
<td>120</td>
<td>1.1°</td>
<td>1.3°</td>
<td>1.7°</td>
<td>2.2°</td>
<td></td>
</tr>
<tr>
<td>240</td>
<td>2.1°</td>
<td>2.7°</td>
<td>3.3°</td>
<td>4.4°</td>
<td></td>
</tr>
<tr>
<td>360</td>
<td>3.1°</td>
<td>4.0°</td>
<td>5.0°</td>
<td>6.5°</td>
<td></td>
</tr>
<tr>
<td>480</td>
<td>4.2°</td>
<td>5.2°</td>
<td>6.5°</td>
<td>8.4°</td>
<td></td>
</tr>
<tr>
<td>600</td>
<td>5.3°</td>
<td>6.4°</td>
<td>8.0°</td>
<td>10.4°</td>
<td></td>
</tr>
</tbody>
</table>

Temperature Change (°C) of 0.200 kg of Water Heated for 300 Seconds

Table D.2

<table>
<thead>
<tr>
<th>Current (amps)</th>
<th>Temperature Change (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1</td>
<td>0.3</td>
</tr>
<tr>
<td>0.2</td>
<td>1.0</td>
</tr>
<tr>
<td>0.3</td>
<td>2.5</td>
</tr>
<tr>
<td>0.4</td>
<td>4.0</td>
</tr>
</tbody>
</table>

Notice that in the course of our experiments we were concerned with four variables (temperature, mass of water, heating time and
heater current), but for the present time we will examine them only two at a time. Finally we will use all of our information to obtain a combined result.

D.2 TEMPERATURE VS. TIME

One relationship that we are interested in is that which may exist between the temperature change in the water and the length of heating time. However, we must be sure to keep the other two variables constant during this search so that our two-variable analysis will be valid. We have already learned that the first step in graphical analysis is the construction of a properly labeled graph. Plot the temperature change (ΔT) against the length of heating time (t) for a mass of 0.200 kilograms of water. Remember to plot the dependent variable (temperature change in this case) on the vertical axis and the independent variable (heating time) on the horizontal axis.

After plotting your points, construct a smooth curve which fits the points as you best estimate. Does your graph appear indicative of any of the proportional relationships that we examined earlier? If so, you may be able to state the proportional relationship that exists between the variables. As we stated earlier, error circles are probably most appropriate here, although students could use error flags if the teacher desires. HPP material to follow uses error flags.

Yes, the direct proportion.
to proceed from a statement of proportionality to an equation it is necessary to determine a constant of proportionality. If you have obtained a straight line, find the constant of proportionality from your graph by determining the slope of the graph. Be sure to indicate units in your result. Now write the equation relating the two variables you have been examining.

You may be curious as to whether or not this same relation applies to heating other masses of water. On the same sheet of graph paper, plot the data for each of the other masses of water. Do all the masses show the same proportional relationship? Do all of the lines have the same equation? What can you conclude?

D.3 TEMPERATURE VS. MASS

Now let's take a look at the relationship that may exist between the change in temperature of the water and the mass of water being heated. The time of heating and current through the heating element are to be maintained at the same values for all the measurements. Plot a graph of temperature change (ΔT) versus mass of water (m) for a heating time of 120 seconds. Does your graph appear to have the
$\Delta T \text{ vs } \frac{1}{\text{M A S S}}$

Temperature Change, $\Delta T$ (°C)

$\frac{1}{\text{M A S S}} \times \frac{1}{m} \left(\frac{1}{\text{kg}}\right)$

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same general shape as one of those already shown? Do we have a technique for finding its slope? If not, we need to try something different before we can proceed with the task of writing an equation.

Your graph probably has a shape resembling the inverse proportion curve discussed earlier. If, in fact, the relationship is a simple inverse proportion $\Delta T = 1/m$, it is relatively easy to show this. Simply plot $\Delta T$ versus $1/m$ and a straight line should be obtained, from which a slope can be found. Try it.

You should now be able to write the equation describing the relationship that exists between temperature change of the water and mass of water for a heating time of 120 seconds.

You may have a suspicion that the same proportional relationship holds true for other masses of water. If you wish, go ahead and graph data for the other masses of water to find the answer for yourself.

D.4 TEMPERATURE VS. CURRENT

Now that we have been reinforced by our success with the first two relationships, let's proceed on to the third. Here we are interested in the relationship that may exist between the temperature change in the water

\[
\Delta T = \frac{\Delta T_{20\text{sec}}}{m} = \frac{k}{m}
\]

$\Delta T_{20\text{sec}} = \frac{1}{m}$ gives a straight line since $\Delta T = \frac{1}{m}$.

$k = 0.33^\circ C\cdot Kg$

$\Delta T_{20\text{sec}} = k \left( \frac{1}{m} \right)$ or $\Delta T = \frac{k}{m}$

$\Delta T_{20\text{sec}} = \frac{.33^\circ C\cdot Kg}{m}$

It does.
being heated (ΔT) and the quantity of current flowing through the heating element (I). Construct your graph, plotting ΔT against I. Something new should be discovered with this graph, since it seems indicative of a power curve.

Before we can even write the correct proportional relationship between the variables, however, we need to know the value of the unknown exponent. There was actually a similar problem in the case of the inverse relation discussed in Section D.3. There we assumed the simplest possibility, viz., a simple inverse of the form y=1/x, and came out all right. In the general case, however, there are several ways to proceed. We could look to the theory of the experiment and make an educated guess as to the power relation. Before the course is over, we will know enough about electricity to be able to do this, but for the present we will not go about it this way. Another approach uses slightly sophisticated mathematical techniques involving logarithms, but this will not be necessary here. The approach we will take, an approach which is used surprisingly often in scientific research, is that of trial and error. This
\[ \Delta T \text{ vs } I \]

- Temperature Change, $\Delta T$ (°C)
- Current, $I$ (Amp)

Graph showing the relationship between temperature change and current.
$\Delta T$ vs $I^2$

Temperature Change, $\Delta T$ (°C)

Current$^2$, $I^2$ (Amp$^2 \times 10^{-2}$)
simply means that we try plotting $\Delta T$ against different powers of $I$ until hopefully we find a power that will give us a straight line. Plotting $\Delta T$ against $I^2$ is probably a good way to start. Try it. If you obtained a straight line you can now write the proportional relationship and equation. If you didn't obtain a straight line, try plotting $\Delta T$ versus $I^3$ or $I^5$, and see if that does the job. Once you have obtained a straight line, you can proceed.

D.5 GRAND FINALE

By now you should have found the relationship between temperature change and time of heating, temperature change and mass of water, and temperature change and current in the heating element. It is entirely possible that in a real situation all three of the independent variables could be varying during the experiment. How then can we determine the relationship between temperature change and each of them at the same time? It's really quite easy. If the temperature change ($\Delta T$) is directly proportional to the time of heating ($t$), the reciprocal of the mass ($\frac{1}{m}$) and the square of the current ($I^2$), then it is directly proportional to the product of the three.

$\Delta T \text{ vs } I^2 \text{ should give a straight line.}$

$\Delta T = I^2$

$\Delta T = k \frac{t}{m}$

$\Delta T = \frac{27^\circ C}{\text{amp}^2} \cdot I^2$.
Writing an equation from this statement of proportionality involves nothing beyond the techniques we have already utilized.

Choose any one of your three completed graphs ($\Delta T$ versus $t$, $\Delta T$ versus $\frac{1}{m}$, or $\Delta T$ versus $l^2$). In each case, note that two of the independent variables were held constant and one varied. Therefore you can read a temperature change corresponding to three known values of the independent variables at any point on the graph. Using this technique, plot a graph of $T$ versus $\frac{t \cdot l^2}{m}$ and write the equation for the line.

E. CONCLUSION

You have now accomplished an important task. The ability to express in the form of an equation the results of laboratory research is the essence of generalization in science. This type of process has led us to many of the best known theories and principles in science, including many of those you will be studying in your first work in physics—concepts of motion.
\[ \Delta T \text{ vs } \frac{I^2 \text{ time}}{\text{mass}} \]

Data from \( \Delta T \) vs time graph using 0.2Kg line for \( \Delta T \) and time

<table>
<thead>
<tr>
<th>( \Delta T )</th>
<th>( \frac{I^2 \text{ time}}{\text{mass}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>192</td>
<td>3.3</td>
</tr>
<tr>
<td>288</td>
<td>5.0</td>
</tr>
<tr>
<td>384</td>
<td>6.5</td>
</tr>
<tr>
<td>480</td>
<td>8.0</td>
</tr>
</tbody>
</table>

*Calculate these values, knowing \( I \), \( t \) and \( m \) for a specific \( \Delta T \).
k value

\[ \Delta T = \text{time} \]
\[ \Delta T = \frac{1}{\text{mass}} \]
\[ \Delta T = l^2 \]
\[ \Delta T = \frac{k \times \text{time} \times l^2}{\text{mass}} \]

Solve for \( k \),

\[ k = \frac{\Delta T \times \text{mass}}{\text{time} \times l^2} \]

From either the data table or one of the graphs, find \( \Delta T \), m, I, and t.

\( \Delta T = 1 \), mass = 0.2Kg, I = 0.2amp, time = 300 sec

\[ k = \frac{1 \times 0.2}{3 \times 4} = 1.7 \times 10^{-2} \, ^\circ\text{C Kg sec-amp}^{-2} \]

\( \Delta T = 2.5 \), mass = 0.2Kg, I = 0.3amp, time = 300 sec.

\[ k = \frac{2.5 \times 0.2}{3 \times 9} = 1.8 \times 10^{-2} \, ^\circ\text{C Kg sec-amp}^{-2} \]

\( \Delta T = 4.0 \), mass = 0.2Kg, I = 0.4amp, time = 300 sec

\[ k = \frac{4.0 \times 0.2}{3 \times 16} = 1.7 \times 10^{-2} \, ^\circ\text{C Kg sec-amp}^{-2} \]

Average \( k = 1.73 \times 10^{-2} \, ^\circ\text{C Kg sec-amp}^{-2} \)
Prologue: THE SCIENTIST AND SCIENTIFIC ACTIVITY  
Unit 1 HP  

Chapter 1: MOTION (KINEMATICS)

A UNIVERSALITY OF MOTION  
Read sections 1.1 and 1.2 HP

B POSITION AS A FUNCTION OF TIME  
Read section 1.3 HP

C AVERAGE SPEED  
Read section 1.4 HP

D GRAPHIC PRESENTATION  
Read section 1.5 HP

D.1 Experiment: UNIFORM MOTION  
(HP Ex 4)

E INSTANTANEOUS SPEED  
Read section 1.7 HP

F ACCELERATION BY COMPARISON  
Read section 1.8 HP

F.1 LOOKING FOR LOGICAL CONSEQUENCES OF GALILEO'S HYPOTHESIS  
(Read sections 2.5 - 2.8 HP)

F.2 Experiment: NON-UNIFORM MOTION  
("A 17th Century Experiment")  
(HP Ex 5)
KEY

Transparency.............T
Film Loop................L
References such as L3, L4, etc. refer to sections in the Handbook, not to an actual film loop
Films.....................F
Reader....................R
Programmed Instruction....P
Demonstration...............D
Experiment..................Ex
Activity....................A

*......Top Priority
x......Recommended
+......Optional
R......Reading Only
E......Easy
Hard...Hard
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<th>TEXT SECTION</th>
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<th>EXPERIMENTS</th>
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<th>OTHER STUDENT ACTIVITIES</th>
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<tr>
<td>Motion of things *</td>
<td>✅</td>
<td>D1 Recognizing simple motion +</td>
<td></td>
<td></td>
<td></td>
<td>Quant</td>
<td>Qual</td>
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<tr>
<td>Motion experiment not quite work *</td>
<td>2 days</td>
<td></td>
<td>T1 Stroboscopic measurements +</td>
<td>Making frictionless pucks + Electronic stroboscope + Magnetic timer +</td>
<td></td>
<td>E</td>
<td>H</td>
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<tr>
<td>Linear motion</td>
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<td></td>
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<td></td>
<td>2*</td>
<td>1*</td>
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<tr>
<td>Polishing motion</td>
<td>✗</td>
<td>D2 Uniform motion using accelerometers and dynamics carts</td>
<td></td>
<td>L8 Analysis x of a hurdle race (Part I)</td>
<td>Discuss #9</td>
<td>4x</td>
<td>5x</td>
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<td>E50's &quot;50&quot; meaning of speed</td>
<td>✗</td>
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<td>Uniform motion</td>
<td>✗</td>
<td>Ex4 Uniform motion</td>
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<td>T2 Graphs of various motions</td>
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<td>Instantaneous</td>
<td>✗</td>
<td>D3 Instantaneous speed using strobe photographs of mass on a spring +</td>
<td></td>
<td>T3 Instantaneous speed + T4 Instantaneous rate of change</td>
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<td>Acceleration</td>
<td>✗</td>
<td>D4 Uniform acceleration using liquid accelerometer</td>
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<td>T1 Stroboscopic measurements + T2 Graph of various motions F1 Straight line kinematics (PSSC)</td>
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<tr>
<td>2.1 - 2.4R</td>
<td>After reading these sections, students may view the film &quot;Galileo--An Age of Reason&quot; which summarizes the material and is the epitome of cross-disciplinary instructional learning</td>
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<td>2.5 Galileo chooses a definition for uniform acceleration</td>
<td></td>
<td></td>
<td>Ex 5A 17th century experiment</td>
<td>T6 Derivation of $d=v_0t+\frac{1}{2}at^2$</td>
<td>SG #13, #14</td>
<td>E</td>
<td>H</td>
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<td>2.6 Galileo could not test his hypothesis directly</td>
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<td>2.7 Looking for consequences of Galileo's hypothesis</td>
<td>Ex 7 Measuring acceleration of gravity</td>
<td></td>
<td>Ex7 Measuring acceleration</td>
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<tr>
<td>2.8 Galileo turns to an indirect test</td>
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Chapter II: NEWTON EXPLAINS MOTION

A  MASS AND FORCE
   Read section 3.1 HP

B  VECTORS
   Read section 3.2 HP

C  NEWTON'S LAW
   Read sections 3.3 - 3.7 HP

C.1  Experiment:  NEWTON'S SECOND LAW
     (HP Ex8)

D  NEWTON'S LAWS
   Read sections 3.8 - 3.11 HP
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<tr>
<td>Roughness</td>
<td>D7 Addition of vectors +</td>
<td>L3 Vector Addition *</td>
<td>P 1, 2, 3 Vectors *</td>
<td>Quant</td>
<td>Qual</td>
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<td>Roughness</td>
<td>D8 Direction of v and a +</td>
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<td>Roughness</td>
<td>D10 Non-commutativity (rotation)</td>
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<td>Explanation</td>
<td>D11 Newton's First Law</td>
<td>Newton's First Law</td>
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<td>Learning</td>
<td>of the laws of motion *</td>
<td>D12 Newton's Second Law</td>
<td>Newton's Second Law</td>
<td>4*</td>
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<td>7x</td>
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<td>Learning</td>
<td>Newton's Second Law</td>
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<td>10+</td>
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<td>Learning</td>
<td>D13 Effects of friction on a</td>
<td>F2 Inertia (PSSC)</td>
<td>Newton's Second Law</td>
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<td>Ex8 Newton's Second Law</td>
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<td>Accelerometer +</td>
<td>17x</td>
<td>15+</td>
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<td>Ex9 Inertial and gravitational mass</td>
<td>D14 Demonstration with rockets + D15 Making an inertial balance</td>
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<td>Learning</td>
<td>3 Days</td>
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<td>Newton's 2nd Law</td>
<td>*</td>
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<td>D16 Action-Reaction (rope) + No. 1</td>
<td></td>
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<td>Qual</td>
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<td>D17 Action-Reaction (rope) + No. 2</td>
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<td>D18 Reaction Forces of a wall +</td>
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<td>T8 The tractor's paradox</td>
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<td>Action-reaction model</td>
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<td>Using Newton's Laws of</td>
<td>2 days</td>
<td>D19 Newton's third law +</td>
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<td>Motion</td>
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<td>D20 Action-reaction (car) +</td>
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<td>D21 Action-reaction (nail) +</td>
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<td>D22 Action-reaction (jumping) +</td>
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<td>Nature's R forces</td>
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**Notes:**
- Quant: Quantitative
- Qual: Qualitative
- T8: The tractor's paradox
- Using Newton's Laws of Motion
- Nature's R forces
Derivations of $v_f = v_i + at$, $d = v_i t + \frac{1}{2}at^2$ and $v_f^2 = 2ad$ (when $v_i=0$)

which are required to solve the problems in Chapter 2 of Harvard Project Physics

Definitions: (1) $\bar{v} = \frac{d}{t}$ (2) $\bar{v} = v_i + v_f$ (3) $a = \frac{\Delta v}{\Delta t}$

(a is assumed to be constant in all cases.)

Derive $v_f = v_i + at$

(4) $a = \frac{v_f - v_i}{\Delta t}$ where $\Delta t = t_2 - t_1$ and $v_f - v_i = \Delta v$

(5) $at = v_f - v_i$ Multiply through by $t$:

(6) $v_f = v_i + at$ Add $v_i$ to both members:

Derive $d = v_i t + \frac{1}{2}at^2$

where $\bar{v}$ (in Eq. 1) = $\bar{v}$ (in Eq. 2)

(7) $\frac{\Delta d}{\Delta t} = v_i + \frac{v_f}{2}$

substitute $v_f = v_i + at$ into equation (7)

(8) $\frac{d}{t} = \left( v_i + at \right) + v_i$ Rearrange:

(9) $\frac{d}{t} = 2v_i + \frac{at}{2}$ Simplify and multiply through by $t$:

(10) $d = v_i t + \frac{1}{2}at^2$

Derive $v_f^2 = 2ad$

Since (6) $v_f = v_i + at$ rearranged gives $t = \frac{v_f - v_i}{a}$

which may be substituted into (7)

(11) $\frac{\Delta d}{\Delta t} = v_i + v_f$ for $t$ Rearrange:

(12) $2ad = v_f^2 - v_i^2$ but when $v_i = 0$ then (13) $v_f^2 = 2ad$
Chapter III: Multi-Dimensional Motion

A  Curved Paths
    Read sections 4.1 - 4.4 HP

A.1 Experiment: Trajectories - I
    (HP Ex 10)

B  Circular Motion
    Read sections 4.5 - 4.7 HP
Solutions to Chapter 4
additional problems

(1) a. \( d = \frac{1}{2} a t^2 \)
   \[ d = \frac{1}{2}(10\text{m/sec}^2)(6\text{ sec})^2 \]
   \[ d = 180\text{ meters} \]

b. \( d = vt \)
   \[ d = 50\text{m/sec} \times 6\text{ sec} \]
   \[ d = 300\text{ meters} \]

(2) a. \( d = vt \)
   \[ d = 50\text{m/sec} \times 2\text{ sec} \]
   \[ d_x = 100\text{ meters} \]

\[ d_y = \frac{1}{2} a t^2 \]
\[ d_y = \frac{1}{2}(10\text{m/sec}^2)(2\text{ sec})^2 \]
\[ d_y = 20\text{ meters} \]

\[ d = \sqrt{(d_x)^2 + (d_y)^2} \]
\[ d = \sqrt{(100\text{m})^2 + (20\text{m})^2} \]
\[ d = \sqrt{10,400} \]
\[ d = 102\text{ meters} \]

b. direction of displacement = \( \frac{20\text{m}}{100\text{m}} \)
   \[ = \frac{1}{5} = .2 = \tan \theta = 11.3^\circ \]

(3) a. \( v_x = 50\text{m/sec} \)
   \[ v_y = at \]
   \[ v_y = 10\text{m/sec}^2 \times 2\text{ sec} \]
   \[ v_y = 20\text{m/sec} \]

b. \( v_z = \sqrt{(v_x)^2 + (v_y)^2} \)
   \[ v_z = \sqrt{(50\text{m/sec})^2 + (20\text{m/sec})^2} \]
   \[ v_z = \sqrt{2900\text{m/sec}^2} \]

\( \tan \theta = \frac{20\text{m/sec}}{50\text{m/sec}} \)
\[ \theta = 21.8^\circ \]

Additional Problems - Chapter 4, HP
Sec. 4.2 and 4.3

(1) A boy standing on a bridge throws a stone horizontally outward. He times the stone, finding it takes 6 seconds to strike the water below.
   a. How high is the bridge?
   b. How far did the stone travel horizontally before striking the water if thrown with an initial speed of 50 m/sec?

(2) a. What was the magnitude of the displacement of the stone in the problem above after 2 seconds?
   b. What is the direction of the displacement? (Express as a ratio of vertical/horizontal distances.)

(3) a. In problem 1, what is the speed of the stone along the x & y axes after 2 seconds?
   b. What is the magnitude of the velocity of the stone after 2 seconds?
   c. What is the direction of the velocity of the stone after 2 seconds? (Express as ratio of vertical/horizontal velocity.)
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<td>4.1 A trip to the moon</td>
<td></td>
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<td>E H E H</td>
<td>E H</td>
<td>E H</td>
<td>1 2*</td>
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<tr>
<td>4.2 Projectile motion</td>
<td>3 Days</td>
<td></td>
<td>T9 Projectile motion</td>
<td></td>
<td>A10</td>
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<td>3* 4*</td>
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<td>Ex10 Trajectories - I</td>
<td></td>
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<td>Generating a cycloid</td>
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<td>4.3 What is the path of a projectile?</td>
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<td>T10 Path of a projectile + Film Free fall and projectile motion(PSSC)*</td>
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<td>4.4 Galilean relativity</td>
<td>1 Day</td>
<td></td>
<td>D23 Frames of reference x D24 Inertial vs. non-inertial reference frames +</td>
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<td>5* 6* 7* 10* 81</td>
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<td>5 Days (cont)</td>
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<td>L4 A Matter of relative motion</td>
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<td>11 F4 Frames of reference (PSSC)</td>
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<td>L5 Galilean relativity 1 (ball dropped from a ship)</td>
<td>E H E H</td>
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<td>L6 Galilean relativity 2 (object from an aircraft)</td>
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<td>L7 Galilean relativity 3 (projectile fired vertically)</td>
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<td>Ex12 Circular Motion</td>
<td>D25 Uniform</td>
<td>A12 Motion in a rotating reference frame</td>
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<td>4.5 Circular motion</td>
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<td>11+ Some centripetal force activities</td>
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<td>19+ 11+</td>
<td>9+ 12+</td>
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<td>4.7 Motion of earth satellites *</td>
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<td>D26: Simple harmonic motion +</td>
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<td>Measuring unknown frequencies +</td>
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<td>Quant E H Qual E H</td>
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<td>T11 Centripetal acceleration (graphical) *</td>
<td>T12 Centripetal acceleration derivation *</td>
<td>F5 Vector kinematics (PSSC) +</td>
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Chapter IV: CONSERVATION

A  CONSERVATION OF MASS
   Read section 9.1 HP

B  COLLISIONS
   Read section 9.2 HP

B.1 Experiment:  COLLISIONS IN ONE
                 DIMENSION
                 (HPEx 22)

B.2 Experiment:  COLLISIONS IN TWO
                 DIMENSIONS
                 (HPEx 23)

C  CONSERVATION OF MOMENTUM
   Read sections 9.3 - 9.7 HP
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<td>1</td>
<td><strong>E23</strong> Collisions in two dimensions</td>
<td><em>F17 Elements, compounds, and mixtures (PSSC)</em>*</td>
<td><strong>Rough Experiments</strong></td>
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<td>4</td>
<td><strong>E22</strong> Collision in one dimension</td>
<td><em>F: The perfection of matter (Nuffield Foundation)</em></td>
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<td>7</td>
<td><strong>T20</strong> Equal mass collision</td>
<td><em>L18 Perfection of one-dimensional collision</em></td>
<td><strong>Other Activities</strong></td>
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<td><strong>T21</strong> Unequal mass collision</td>
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<td><em>L20 Interesting case of elastic impact</em></td>
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**Quantitative Observation**

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**Qualitative Observation**

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<td>L24 Scattering of a cluster of objects</td>
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<td>L21 Two-dimensional collisions</td>
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<td>L22 Further examples of two-dimensional collisions</td>
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Chapter V: ENERGY - WORK

A WORK AND KINETIC ENERGY
   Read section 10.1 HP

B POTENTIAL ENERGY
   Read section 10.2 HP

C CONSERVATION OF MECHANICAL ENERGY
   Read section 10.3 HP

D FORCES THAT DO NO WORK
   Read section 10.4 HP

E ENERGY IN BIOLOGICAL SYSTEMS
   Read section 10.8 HP

F CONSERVATION OF ENERGY
   Read section 10.9 HP
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<td>A method of measuring energy—nails driven into wood</td>
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<td>L34: Range of two-dimensional elastic collisions</td>
<td>L32: Gravitational potential energy</td>
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<td>L35: Conservation of energy</td>
<td>F20: Energy and work (PSSC)</td>
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<td>L26: Speed of rifle bullet Method I</td>
<td>L33: Kinetic energy</td>
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<td>L27: Speed of rifle bullet Method II</td>
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<td>L10: Air-craft take-off</td>
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<td>10.8 Energy in biological system</td>
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<td>One student ?</td>
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<td>10.9 The law of conservation of energy</td>
<td>2 Days</td>
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<td>L29 Colliding freight cars x L30 Dynamics of a billiard ball</td>
<td>Rubberband engines</td>
<td>L29 Colliding freight cars x L30 Dynamics of a billiard ball</td>
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<td>L4 A matter of relative motion + F21 Conservation of energy(PSSC)</td>
<td>Energy analysis of pendulum swing</td>
<td>L4 A matter of relative motion + F21 Conservation of energy(PSSC)</td>
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<td>L4 A matter of relative motion + F21 Conservation of energy(PSSC)</td>
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Chapter VI: KINETIC THEORY OF GASES

A  KINETIC THEORY
Read sections 11.1 - 11.5 HP

A.1 Experiment: BEHAVIOR OF GASES
(HPEx 29)

B  SECOND LAW OF THERMODYNAMICS
Read sections 11.6 - 11.8 HP
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<td>11.1 Heat and mechanical energy</td>
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<td>F22 Mechanical energy and thermal energy (PSSC) x</td>
<td>Scirling hot air engine + Equivalent of heat +</td>
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<td>11.2 A model for the gaseous state</td>
<td>4 days</td>
<td>Ex 29 Behavior of gases +</td>
<td>D35: Diffusion of gases +</td>
<td>F23 Demonstrating gas laws (Coronet) x</td>
<td>F24 Gas laws and their applications (EBF) x</td>
<td>Gas Laws: How the amateur can experiment with films only one molecule thick x</td>
<td>1* 2* 3+ 4* 5+ 10x</td>
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<td>11.3 The speeds of molecules</td>
<td>*</td>
<td>Ex 28 Monte Carlo experiment on molecular collisions *</td>
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<td>L36 Kinetic theory - Gas Model x</td>
<td>Molecular speed distribution +</td>
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<td>R8: The Law +</td>
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<td>11.4 The sizes of molecules</td>
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<td>F25 Molecular theory of matter (EBF) x</td>
<td>Wie Gross ist ein Molekul?</td>
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#PSSC Ex on molecular layers option here.
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<td>Studying the properties of gases experimentally</td>
<td>13* 8*</td>
<td>R5 The Great molecular theory of gases R6 On the kinetic theory of gases</td>
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<td>11.6 The second law of thermodynamics and the dissipation of energy</td>
<td>1 Day</td>
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<td>Rockets</td>
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<td>15* 17*</td>
<td>R4 The barometer story</td>
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<td>11.7 Maxwell's demon and the statistical view of the second law of thermodynamics</td>
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<td>Perpetual motion mechanics</td>
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<td>R1 The law of disorder R10 Maxwell R11 Maxwell's demon</td>
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<td>2 Days</td>
<td>L37 Reversibility of time</td>
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<td>Problems of scientific and technological growth</td>
<td>19* 20* 21*</td>
<td>R9 The Arrow of time R12 Randomness of the 20th century</td>
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CHEMICAL REACTIONS
Chapter 1: THE MOLE AS A COUNTING UNIT

A COUNTING THINGS
A.1 Exercise: COUNTING "BB'S"

B COUNTING UNITS

C THE MOLE

D ATOMIC WEIGHT
D.1 Experiment: COPPER WIRE, SILVER NITRATE AND MOLES
D.2 Experiment: FROM MOLES OF SILVER TO MOLES OF SILVER CHLORIDE

E FORMULAS FOR MOLECULES

F MOLECULAR WEIGHTS

G RELATIVE WEIGHTS OF MOLECULES
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<td>Exercise: Counting &quot;BB's&quot;</td>
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<td>Formulas for molecules</td>
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9, 10, 11, 12, 13

14, 15, 16, 17
A. COUNTING THINGS

In previous chapters a model for the gaseous state has been examined. Speeds and sizes of molecules were discussed and the ideal gas law was derived by considering topics such as pressure, momentum and temperature of gases.

\[ P = kDT \]

If man wants to communicate about particles or things, he must be able to describe them. What is the shape of a gas particle? Is it spherical? How many are there in a given volume? How does one count these extremely small particles that go dashing randomly about in a container? Let's first think about counting these gas particles. How does man usually count things and what are some alternative ways of counting things?

A.1 Exercise: COUNTING "BB'S"

Although "BB's" are enormously large when compared to gas particles, the practice in devising an alternative method of counting these fairly small particles will be helpful.

Count out ten BB's; don't laugh--do it. How did you count this small number of parti-
Students may decide on other "easy" ways of counting BB's. If they decide--for example--on volume, you might ask them, "What could possibly make your counting by volume method inaccurate?" Does massing or volumetric measurement give you the better results?

The students who first found the mass of ten BB's, set their balance at one hundred times the mass of ten BB's and then poured BB's onto the balance until it "tripped," are proceeding in the right direction.

Now count out one thousand BB's. Think about alternative ways of counting. What is the easiest way to count one thousand BB's? Do it. What would be the easiest way to count all the BB's in the classroom?

Then counting particles is an easy task. All that is needed is the mass of a number of particles. But, you say, "ten gas particles mass so little (about $10^{-21}$ grams) that our laboratory balances cannot mass a unit of ten."

What we really need is an appropriate counting unit for gas particles.

B. COUNTING UNITS

One of the most used counting units is the dozen. We can think of the mass of twelve cows or people, but the mass of twelve atoms is very little different from the mass of ten. Obviously the dozen is not a good counting unit to use when working with gas particles. What about the gross (144 things)? No, since a gross of gas particles masses about $10^{-21}$ grams. We need a "really big" counting unit.

C. THE MOLE

Such a unit already exists. It is called the mole. It has nothing to do with little animals that live in the ground, unless you wish to count them. The mole is $6.02 \times 10^{23}$ things,
just as the dozen is 12 things. One can talk about a mole of cows or mosquitoes; however the use of such a large counting unit for this purpose is ridiculous. One mole of cows would mass about $10^{26}$ Kg! This is approximately sixteen times greater than the mass of the earth! Gas particles, however, have such small masses that a mole of gas can be massed on a typical laboratory balance.

D. ATOMIC WEIGHT

If our gas particle happens to be monoatomic (composed of just one atom), then the weight in grams of one mole of the gas is its atomic weight. The weight of one mole of helium gas, for example, is four grams.

Find a table of atomic weights and determine the weight of one mole of neon, krypton and xenon. The idea of atomic weight does not stop with just gases. The weight in grams of one mole of any mono-atomic element is the atomic weight of that element. Notice the term 'weight' is used in the definition of ATOMIC WEIGHT. We know the definition should read, "the mass in grams of one mole of atoms is the atomic weight." Traditionally chemists have used the term weight when they mean mass. Although tradition can be confusing, most

The weight of one mole of
Neon = 20.2 grams
Krypton = 83.8 grams
Xenon = 131.3 grams

You may have noticed the change from the term "mass" to the term "weight." "Physically" speaking, mass is the proper term, but the chemistry that follows uses the term "weight." Atomic wt., molecular wt. and hence the numbers used on periodic tables are relative wts., based on carbon twelve. The numerical distinction between the mass of a mole of atoms and the weight (relative) of a mole of atoms is negligible when working with ordinary massing techniques. Prolonged discussion of the topic mass vs. weight is inappropriate at this time because:

1. Confusion usually results.
2. The time used could be better spent on more significant topics.

However, if your students open the topic - have at it.

See Chem Study T. G., Experiment #7, for details concerning materials, etc.

Purified water should be used in this experiment. Small laboratory de-ionizer units, such as the Crystalab Deeminizer are very useful. They operate by ion exchange similarly to a home water-softener unit. Distilled water is, of course, satisfactory.

\[ Cu + 2AgNO_3 \rightarrow Ag + Cu(NO_3)_2 \]

areas of study are bound by it; and we must simply remember the chemist usually means mass when he says "weight."

D.1 Experiment: COPPER WIRE, SILVER NITRATE AND MOLES

In this experiment you will prepare a solution of silver nitrate. You will also weigh a length of copper wire, and after it has remained in the silver nitrate solution over night you will weigh it again. In this and the following experiment you will count, by weighing, the number of moles of particles involved.

Copper wire is composed of copper particles. It is an element; it contains only one kind of atom. The symbol for copper is Cu.

Silver nitrate is a compound which has the formula \( AgNO_3 \). You will find your laboratory work even more rewarding if you prepare a data table before you come to lab.

Your data should include the following:

- Weight of Cu wire before immersion
- Weight of Cu wire after immersion
- Weight change of Cu wire
- Weight of vial plus AgNO_3
- Weight of vial
- Weight of AgNO_3
- Weight of beaker plus residue
Weight of beaker
Weight of residue

Obtain a 30 cm piece of copper wire. Clean it with steel wool. Then form a coil by wrapping the wire around a small test tube. Leave about 5 cm for a handle.

Carefully weigh the copper wire. Also weigh a clean dry 250 ml beaker. Weigh the vial of silver nitrate provided by the instructor. Do not use tap water because of impurities. Fill the beaker about two-fifths full with de-ionized water. Add the silver nitrate (AgNO₃) to the water. Stir gently with a glass rod until all the silver nitrate (AgNO₃) crystals have dissolved. Weigh the empty vial. Silver nitrate (AgNO₃) solution or solid will stain your skin and clothing. It will be easy for your instructor to determine who is the sloppy experimenter.

Now place the copper wire in the beaker with the coil immersed in the AgNO₃ solution. Observe and record any changes that take place for at least five minutes. Then place the beaker, covered with a watch glass, in the designated place until the next laboratory period.

When you return to the laboratory, be
very careful not to disrupt the contents of the beaker until you have had a chance to observe the contents. Record your observations.

Shake the crystals off the coil and lift the coil out of the solution. Use a wash bottle to rinse into the beaker any crystals which tend to adhere to the coil. Now set the coil aside to dry after you have rinsed it in acetone. Weigh the coil when it is dry.

Let the crystals settle in the beaker and carefully decant the solution. Decant means to pour off the liquid and leave the solid behind.

See Figure D.2.A and B

Next add 5 ml of dilute AgNO₃ solution and stir gently until any Cu flecks left in the residue in the original beaker disappear. Carefully decant again. Wash the residue with 10 ml of water and carefully decant again. You may neglect the few particles which may float over with the wash water because their weight is usually insignificant. (Uncertainty in measurement?)

Now the residue must be dried. Your instructor will suggest a suitable method. This usually takes less than 24 hours.

When you return to the laboratory allow the beaker and its contents to cool before
weighing. Should you use the same balance you used previously? Record the weight.

How can you be certain the contents are dry? COVER THE BEAKER AND CONTENTS WITH A WATCHGLASS AND SAVE IT FOR THE NEXT EXPERIMENT.

Calculations:

1. Calculate the part of a mole of copper that went from the wire into the solution.
2. The residue is silver (Ag). Calculate the part of a mole of silver you obtained.
3. What is the ratio of moles of silver to moles of copper? Express your answer using scientific notation.
4. What you have observed can be partially described by the following:
   \[ \text{1 mole of copper (solid)} + \underline{\text{mole(s) of silver nitrate (in water)}} \rightarrow \underline{\text{mole(s) of silver (solid)}} + \underline{\text{mole(s) of copper nitrate (in water).}} \]
   Using your results, write the proper whole numbers in the blanks.
5. How many atoms of Cu were lost by the wire?

Students should use the same balance. The following example might help students see why:

Suppose you wish to know the mass of your pet cat. An easy way to find it would be to first mass yourself on the bathroom scales and then mass yourself holding the cat. The difference in mass would be the mass of the cat, as long as you used the same scale. It would not matter if the scale had a negative or positive instrumental error. However, if you used two balances, one of which had a negative error, you might think your cat has lost or gained mass at a fantastic rate, depending on which balance you first use. Why?

Students can be certain the contents are dry by alternately heating and weighing the beakers. If the mass does not change, then the contents are dry.

Answers to Calculations

1. weight change of Cu wire = weight of Cu/mole Cu
   \[ \text{weight change of Cu wire = wt. of Cu/mole Cu} \]
   moles Cu

2. weight of residue = moles wt. of Ag/mole Ag of Ag

3. moles of Ag should, with moles of Cu in the range of uncertainty, equal 2:1

4. \[ +2 \rightarrow 2 + 1 \]

5. \[ \text{(moles)}(6.02 \times 10^{23} \text{ atoms}) = a \]
   \[ \text{(1)}( \frac{\text{mole}}{1}) \]
6. \((\text{moles})(6.02 \times 10^{23}\text{atoms}) = a\) (mole)

7. Same answer as calculation #3.

8. The blue color is due to the aquated copper (II) ion.

6. How many atoms of silver were left as the residue?

7. What is the ratio of atoms of silver to atoms of copper (from your calculations in 5 and 6)?

8. Your instructor or you may want to do a frequency distribution for the results of this experiment. (How often did students get the same answer for the ratio of copper to silver?)

If you do the frequency distribution, please consider how the range of values obtained compares with the uncertainty you consider justifiable from your measurements.

What do you think causes the blue color of the solution?

D.2 Experiment: FROM MOLES OF SILVER TO MOLES OF SILVER CHLORIDE

In your next experiment the silver produced in the previous experiment is used to form a water solution of AgNO₃. This is done by allowing the silver to react with nitric acid \((\text{HNO}_3)\). A water solution of sodium chloride \((\text{NaCl})\) is then added to the silver nitrate solution and silver chloride is produced.

Many techniques are reviewed in this ex-
periment and a few new ones are learned. Learn as much as you can of these techniques—careful weighing, decanting, washing, drying and filtering—because they will be used often in future experiments.

Remember to record your data carefully. Include the uncertainty and units in each measurement. You should have a data table before you come to the laboratory.

Your data should include the following:

- Weight of silver from previous experiment
- Weight of silver nitrate used in previous experiment
- Weight of beaker #1
- Weight of beaker #1 and solid AgNO₃
- Weight of beaker #1, filter paper, and solid AgCl
- Weight of beaker #2
- Weight of beaker #2 and solid NaCl
- Weight of filter paper
- Weight of beaker #2 and solid residue

Record the weight of the beaker used in the last experiment—the one with the solid silver in it. Label this beaker #1. Also record the weight of silver nitrate used and the weight of silver produced.
DO THIS PART IN THE HOOD. NO IS TOXIC.

By using two different size beakers there is less chance that the student will confuse the two solutions.

Add 15 ml of nitric acid (HNO₃), labeled 6 M HNO₃, to the beaker containing the silver. Don't inhale any of the poisonous reddish-brown fumes of nitrogen dioxide (NO₂) which form as the silver dissolves. Put an identifying mark on your beaker (#1) and leave it in the fume hood or near an open window to dry for about 24 hours.

When you return to the laboratory, weigh the beaker which now contains AgNO₃. (Does this material stain clothing?)

Add 15 ml of de-ionized water to the AgNO₃ in beaker #1. Stir until no more change takes place.

Pour out about 1/2 teaspoon of sodium chloride (NaCl) as shown below:

- Rotate the bottle to obtain the approximate amount of solid

Now label a clean dry 100 ml beaker as #2 and weigh it to the nearest 0.01 g.

Adjust your balance so it reads about 2.0 to 2.5 grams greater than beaker #2. Carefully add the NaCl from the paper to beaker #2 until the balance "trips." Then carefully weigh the beaker with the NaCl and record this weight on your data table.

Add about 15 ml of de-ionized'water to the
NaCl in beaker #2. Stir until no more change is observed.

Add the NaCl solution slowly to the AgNO₃ solution while rapidly stirring the AgNO₃ solution. The white solid produced is the compound silver chloride (AgCl). Rinse beaker #2 with about 5 ml of de-ionized water.

![Rinsing a Beaker with a Wash Bottle](image)

**Figure D.4**

Add the rinse water to the mixture in beaker #1. Rinse beaker #2 again with de-ionized water and discard the rinse water. Beaker #2 will be used later.

Now heat beaker #1 to boiling for about 2 minutes. The solution should become reasonably clear as the precipitate settles. If you place a stirring rod in the beaker, it will help prevent unsteady boiling (bumping).

Weigh a piece of filter paper. Then fold it as shown below.
Place beaker #2 under the funnel to catch the filtrate (the liquid that comes through the filter paper). Decant the liquid from beaker #1 into the funnel. A small amount of precipitate may go into the filter paper, but try to keep most of it in beaker #1 for washing.

Wash the precipitate in beaker #1 with about 15 ml of de-ionized water, stirring with a glass rod to aid the washing. Decant the wash water into the funnel. Repeat the washing procedure and again decant the wash water into the funnel.

After the filtration is complete place the filter paper, partially opened, in beaker #1.

Now place both beakers, #1 and #2, in a
place designated by your teacher for drying. **Be certain** you can recognize your beakers.

When you return to the laboratory, weigh both dry samples and record the weights. (How can you be certain the samples are dry?) Save the silver chloride as directed by your teacher.

**Calculations:**

Fill in the following table:

<table>
<thead>
<tr>
<th></th>
<th>Wt.(gms.)</th>
<th>Moles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ag (previous exp.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AgNO₃ (previous exp.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AgNO₃ produced this exp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NaCl added</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AgCl (in beaker #1)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. How does the number of particles of AgNO₃ produced in this experiment compare with the number of particles of AgNO₃ used in the previous experiment? How do you account for any similarity or difference?

2. Compare the sum of the weights of the AgNO₃ and NaCl used with the sum of the weights of the AgCl and the residue in beaker #2. What is the significance of these results? Per-

Students find moles by dividing weight by weight/mole.

1. The moles of AgNO₃ produced should be equal to the moles of AgNO₃ used in the previous experiment. Conservation of Mass (Ag atoms) should account for the similarity and the uncertainty associated with measurement should account for any difference.

2. Again, mass is conserved in chemical reactions. The fact that all of the reactant, NaCl, might not have been used does not invalidate the results.
3. The moles of silver remain constant.

4. The color is due to impurities, probably from inadequate washing.

Have a few copies of The Handbook of Chemistry and Physics on hand.

haps some class data would be helpful.

3. Check on the moles of silver atoms involved in both experiments. Considering the uncertainty associated with your measurements, what can you conclude about the number of moles of silver atoms involved in this series of chemical changes?

4. Pure AgNO₃ is a white solid. How do you account for any color which may be present in the prepared samples?

If you have time, you may wish to devise an experiment to determine the composition of the residue in beaker #2. Check with your teacher.

E. FORMULAS FOR MOLECULES

Molecules are made of atoms in definite arrangements and definite numbers. Symbolism aids us in demonstrating the number of kinds of atoms in a molecule. For example, H₂ means two atoms of hydrogen make up a molecule of hydrogen gas. A molecule of water is composed of two atoms of hydrogen and one atom of oxygen. The formula for water, H₂O, conveys all this information. The molecular formula for ammonia is NH₃. What does the formula represent? It is possible to find tables of the symbols commonly used for
elements. The Handbook of Chemistry and Physics is a good place to look for molecular formulas.

It is often desirable to translate molecular formulas from digits to words. This is done by using the appropriate Greek prefix from the list below:

- **mono**... one
- **di**... two
- **tri**... three
- **tetra**... four
- **penta**... five
- **hexa**... six
- **hepta**... seven
- **octa**... eight
- **nona**... nine
- **deca**... ten

Often no prefix is used and in these cases 'mono' is understood.

Examples:

<table>
<thead>
<tr>
<th>Molecular formula</th>
<th>translation in words</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂</td>
<td>carbon dioxide</td>
</tr>
<tr>
<td>PBr₃</td>
<td>phosphorus tribromide</td>
</tr>
<tr>
<td>CCl₄</td>
<td>carbon tetrachloride</td>
</tr>
<tr>
<td>PCl₅</td>
<td>phosphorus pentachloride</td>
</tr>
<tr>
<td>N₂O₅</td>
<td>dinitrogen pentoxide</td>
</tr>
<tr>
<td>NaCl</td>
<td>sodium chloride</td>
</tr>
</tbody>
</table>

F. MOLECULAR WEIGHTS

If a mole of atoms has a definite weight, then a mole of molecules which is composed of atoms in definite numbers must also have a definite weight. One mole of H₂O must contain two moles of hydrogen atoms and one mole of
oxygen atoms. Therefore the molecular weight of \( \text{H}_2\text{O} \) is \( 1 + 1 + 16 = 18 \). The gram molecular weight is the weight in grams of a mole of molecules. What is the gram molecular weight of \( \text{H}_2 \) of \( \text{NH}_3 \)?

G. RELATIVE WEIGHTS OF MOLECULES

Imagine that in your left hand you have a dozen identical red marbles that have a total weight of thirty-two grams and in your right hand a dozen identical white marbles that have a total weight of sixteen grams. The ratio of the total weight in your left hand to the total weight in your right is thirty-two to sixteen or two to one. But--can we determine the ratio of weight of one red to one white marble from this same information? Of course; the ratio of total weights is the same as the ratio of individual weights. Therefore, if the ratio of molecular weights is known, the ratio of weights of individual molecules is also known, if we know there is the same number of each. For example, one mole of oxygen weighs 32 grams and one mole of hydrogen weighs two grams. The ratio of their molecular weights is 32 to 2 or 16 to 1. The ratio by weight of an individual oxygen molecule (\( \text{O}_2 \)) to an individual hydrogen molecule
(H₂) must also be 16 to 1. This comparison of the weight of a mole of one substance to the weight of a mole of a second substance yields a meaningful ratio because a mole of any substance contains the same number of molecules \((6.02 \times 10^{23})\) as a mole of any other substance.
Exercises for Home, Desk and Lab (HDL)

1. Ten moles of stone would approximate the mass of the earth.

2. Each inhabitant would receive $2 \times 10^{14}$ dollars/individual.

3. Gold weighs $3.28 \times 10^{-2}$ g/atom.

4. There would be $6 \times 10^{13}$ silver atoms in $10^{-8}$ grams.

5. a. $\frac{1}{3}$ mole
   b. 0.05 mole

6. Potassium, calcium, cobalt, lead, scandium.

7. a. Si $O_2$
    b. S $Cl_2$
    c. $NF_3$
    d. Al $Br_3$
    e. $N_2I_2$

8. A stone about the size of a grapefruit weighs roughly 1 kilogram. How many moles of such stones would be needed to account for the entire mass of the earth, which is about $6 \times 10^{24}$ kg?

9. If we had one mole of dollars to divide among all the people in the world, how many dollars would each of the three billion inhabitants receive?

10. The gram atomic weight of gold is 197. Determine the weight in grams of one gold atom.

11. The most delicate balance can detect a change of about $10^{-8}$ grams. How many atoms of silver would be in a sample of that weight? The gram atomic weight of silver is 108.

12. How many moles of atoms are in a 9.0 grams of aluminum or 0.83 grams of iron?

13. What do the following symbols represent? K, Ca, Co, Pb, Sc.

14. Write formulas for:
   a. silicon dioxide
b. sulfur dichloride  
c. nitrogen trifluoride  
d. aluminum tribromide  
e. dinitrogen diiodide

(8) a. What does the molecular formula $\text{CCl}_4$ mean?  
b. What information is added by the following structural formula?  
Guess!  
\[ \text{Cl} \]  
\[ \text{Cl} - \text{C} - \text{Cl} \]  
\[ \text{Cl} \]

(9) How many moles of oxygen atoms are in one mole of perchloric acid molecules ($\text{HClO}_4$)?

(10) Consider the following data:

<table>
<thead>
<tr>
<th>Element</th>
<th>Atomic Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>12.0</td>
</tr>
<tr>
<td>Y</td>
<td>19.0</td>
</tr>
</tbody>
</table>

X and Y combine to form a new substance Z. If one mole of X combines with four moles of Y atoms to give one mole of Z,  
a. What is the formula of Z in terms of X and Y?  
b. What is the weight of one mole of Z?

(11) A chemist weighs out 10.0 grams of carbon dioxide, 10.0 grams of water and 10.0 grams of magnesium. How
(12) The molecular formulas and weight of one mole of each of the following is:

- Carbon monoxide (CO) 28.0 g
- Silicon dioxide (SiO₂) 60.1 g
- Phosphorus tribromide (PBr₃) 271 g
- Titanium tetrachloride (TiCl₄) 190 g
- Uranium hexafluoride (UF₆) 352 g

(13) Calculate the molecular weight of each of the following:

a. SiF₄
b. HF
c. F₂
d. Xe
e. NO₂

(14) The ratio of the weight of a liter of chlorine gas (Cl₂) to the weight of a liter of oxygen gas (O₂) is 2.22 (both measured at room temperature and pressure). Calculate the molecular weight of chlorine.

(15) Gaseous uranium hexafluoride is important in the preparation of uranium as a source of atomic energy. A flask is filled with gaseous uranium hexafluoride. The uranium hexafluoride, UF₆, is found to weigh 7.04 grams. The same many moles of each substance does he have?

(12) Write the molecular formulas for the following compounds and give the weight of one mole of each: carbon monoxide, silicon dioxide, phosphorus tribromide, titanium tetrachloride, and uranium hexafluoride.
flask is filled with oxygen and, under similar conditions, the oxygen weighs 0.64 grams. What is the ratio of the weight of one uranium hexafluoride molecule to the weight of one oxygen molecule?

(16) Determine the weight ratio of an oxygen atom to a carbon atom.

(17) Oxygen (O₂) is found to weigh 8 times as much as an unknown gas. What is the molecular weight of the unknown gas?

(16) \( O_2 = 16 \text{ g} \)
\( C = 12 \text{ g} \)
\[ \text{ratio} = \frac{16}{12} \]
\[ \text{or weight ratio} = \frac{1.33}{0} \frac{O}{C} \]

(17) Weight ratio \( 8 \frac{g \text{ O}}{1 \text{ g Unknown}} \)

\[ M. \text{W.} \ O_2 = 32 \text{ g} \]
\[ 32 \text{ g} \ O_2 \times \frac{1 \text{ g Unknown}}{8 \text{ g} \ O_2} \]
\[ = 4 \text{ g Unknown gas} \]
Chapter II: COMBINATIONS OF GASES

A  AVOGADRO'S HYPOTHESIS

A.1 Experiment: THE WEIGHTS OF EQUAL VOLUMES OF GASES AT THE SAME TEMPERATURE AND PRESSURE

A.2 THE EFFECT OF BUOYANCY ON WEIGHING

B  THE VOLUME OF A MOLE OF GAS

C  STANDARD TEMPERATURE AND PRESSURE (S.T.P.)

D  TOTAL PRESSURE AND PARTIAL PRESSURE

D.1 Experiment: A QUANTITATIVE INVESTIGATION OF THE REACTION OF MAGNESIUM WITH HYDROCHLORIC ACID
<table>
<thead>
<tr>
<th>SECTION</th>
<th>ROUGH TIME ESTIMATES</th>
<th>EXPERIMENTS</th>
<th>DEMONSTRATIONS</th>
<th>TEACHING AIDS</th>
<th>OTHER STUDENT ACTIVITIES</th>
<th>PROBLEMS</th>
<th>OUTSIDE READING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Combina- of gases Avogadro's thesis</td>
<td>2 Days</td>
<td>The weights of equal volumes of gases at the same temp. and pressure</td>
<td>Film - gases and how they combine CHEMS</td>
<td></td>
<td>1, 2, 3, 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The volume mole of</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5, 6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S.T.P.</td>
<td></td>
<td>A Quantitative Investigation of the Reaction of Mg with HCl</td>
<td></td>
<td></td>
<td>7, 8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total pressure</td>
<td>2 Days</td>
<td></td>
<td></td>
<td></td>
<td>9, 10</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Amadeo Avogadro, an Italian scientist who lived during the 18th and 19th century, noticed that gases combine in small whole number ratios. For example:

Two liters of the gas nitric oxide combines with one liter of the gas oxygen to form \( \frac{5}{6} \) two liters of the gas nitrogen dioxide

One liter of the gas oxygen combines with two liters of the gas hydrogen to form two liters of the gas water.

These combinations in small whole numbers occurred only if the volumes were measured at the same temperature and pressure. What do these observations have to do with particles and counting them?

A. **Avogadro's Hypothesis**

Equal volumes of gases, measured at the same temperature and pressure, contain equal numbers of molecules.

This hypothesis, first made in 1811, is still accepted today. That equal volumes contain equal numbers of particles at similar conditions allows the direct determination of relative weights of gases.
A.1 Experiment: THE WEIGHTS OF EQUAL VOLUMES OF GASES AT THE SAME TEMPERATURE AND PRESSURE

In this experiment you will weigh equal volumes of carbon dioxide and oxygen and perhaps another gas. As you know, the volume of a gas varies with temperature and pressure. Precautions must be taken to keep these variables the same as the gases are weighed. Precautions must also be taken in weighing, because the weight of the gas samples will be at most a few grams. With such small weights, the uncertainty of measurement could destroy the significance of your data. A few greasy finger prints or a moisture droplet are enough to destroy your results. In other words "cleanliness is next to good results."

Before coming to the laboratory read the experiment (pages 67-71) and organize a data table.

Part I. **Weighing carbon dioxide gas \((\text{CO}_2)\)\text{(g)}**

Obtain the apparatus pictured below from your teacher and put it together as pictured.
Figure A. 1

Be certain the large end of the stopper is inside the bag. If you fold the bag around the stopper in small pleats, there will be less chance of leakage. Be careful when you force the medicine dropper into the stopper. Experience shows it is less painful if there is toweling between glass and your hand if the glass breaks. Ask your teacher for some help if you do not already know the proper technique for handling glass tubing.

Press out any air in the bag by smoothing it flat. Then replace the rubber cap on the medicine dropper and weigh the complete apparatus. Don't forget to record the uncertainty.

Remove the rubber cap and connect the apparatus to the $\text{CO}_2(g)$ source supplied by your teacher. Allow the bag to be fully inflated.

Save the stopper assembly for future years.
Hold the apparatus by the stopper and disconnect the rubber tubing from the medicine dropper. Don’t squeeze the bag but let the excess gas escape so that the bag will be at room pressure. Now replace the rubber cap while still holding the bag.

Weigh the bag apparatus with the \( \text{CO}_2 \) gas and record the uncertainty in the measurement. You may wish to repeat the weighing of the \( \text{CO}_2(\text{g}) \) in order to check your work.

Part II. Weighing oxygen gas (\( \text{O}_2(\text{g}) \))

Empty the bag and stopper apparatus. Fill it with \( \text{O}_2(\text{g}) \) using the source supplied by your teacher. Remember to allow the excess \( \text{O}_2(\text{g}) \) out of the bag. Weigh the bag apparatus with the \( \text{O}_2(\text{g}) \) and record with the uncertainty. Remember--the idea is to have equal volumes of gases at the same temperature and pressure. Save the apparatus (bag filled with oxygen) for the determination of volume that follows.

If another gas is available and you have time, you may want to obtain the weight of the third bag of gas.

Part III. Volume determination

Measure the volume of oxygen in the bag.
by the method shown in the picture on the follow-
ing page. (Figure A.2) Completely fill a large bottle or jug (about 2 liters) with tap water. Place a glass square over the mouth of the jug and invert it over the trough of water. Remove the square of glass under water.

Shove one end of a suitable piece of rubber tubing into the jug. Push it up within one centimeter of the bottom of the jug.

Now remove the cap from the medicine dropper and attach it to the other end of the rubber tubing. Gently press on the bag so the gas will displace the water in the bottle. Finally smooth out the bag to remove all the gas.

Quickly pinch the tubing so that the back pressure will not force oxygen (or water if the end of the tube is below the water level) back into the bag. Pull the tube out of the jug and set the jug upright on the table.

Measure the volume of water needed to refill the bottle. Record the volume of the bag. What is the uncertainty of the measurement? Why is the volume needed to refill the bottle the volume of the bag full of gas? Record room temperature and pressure. If you haven't read a barometer before, ask your teacher or someone who has to show you how.
Measuring the volume of a plastic bag

Figure A. 2.
A.2 THE EFFECT OF BUOYANCY ON WEIGHTINGS

Because we live in a sea of air, we usually do not consider its effects on measurements. When any object is weighed in air, it is buoyed up by a force equal to the weight of the volume of air it displaces. When you step on your bathroom scales, you read a weight that is less than your true weight. Your volume is about 70 liters. At normal conditions of temperature and pressure seventy liters of air is about 84 grams or three ounces.

\[(70 \text{ liters}) (1.2 \text{ g/liter}) = 84 \text{ grams.}\]

Unless you have an unusual bathroom scale, the uncertainty of the instrument is much greater than three ounces. We usually neglect the buoyancy effect because it is relatively small.

Suppose that instead of the air around us we were immersed in water and we had the proper scuba equipment. As you probably know, objects are easier to lift when they are submerged in water. The water buoys up a submerged object with a force equal to the weight of the volume of water it displaces. For example, an object with a volume of one liter (1000 ml) when submerged in water displaces one liter of water. Therefore, it is buoyed up with a force of 1000 g, the weight of a liter of water. The same object in air is buoyed up with a
force of about 1.2 grams (the weight of the liter of air it displaces).

In this experiment the apparent weight of the gas is not the true weight of the gas. We must add the weight of the volume of air displaced to the apparent weight of the gas in order to find the true weight of the gas.

The weight of the volume of air displaced can be found by using the following table.

WEIGHT OF A LITER OF AIR IN GRAMS PER LITER, ± 0.01 g, AT VARIOUS TEMPERATURES AND PRESSURES

<table>
<thead>
<tr>
<th>Pressure (mm Hg)</th>
<th>15°C</th>
<th>20°C</th>
<th>25°C</th>
<th>30°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>700</td>
<td>1.13</td>
<td>1.11</td>
<td>1.09</td>
<td>1.07</td>
</tr>
<tr>
<td>710</td>
<td>1.14</td>
<td>1.12</td>
<td>1.10</td>
<td>1.09</td>
</tr>
<tr>
<td>720</td>
<td>1.16</td>
<td>1.14</td>
<td>1.12</td>
<td>1.10</td>
</tr>
<tr>
<td>730</td>
<td>1.18</td>
<td>1.16</td>
<td>1.14</td>
<td>1.12</td>
</tr>
<tr>
<td>740</td>
<td>1.19</td>
<td>1.17</td>
<td>1.15</td>
<td>1.13</td>
</tr>
<tr>
<td>750</td>
<td>1.21</td>
<td>1.19</td>
<td>1.17</td>
<td>1.15</td>
</tr>
<tr>
<td>760</td>
<td>1.23</td>
<td>1.21</td>
<td>1.19</td>
<td>1.16</td>
</tr>
<tr>
<td>770</td>
<td>1.24</td>
<td>1.22</td>
<td>1.20</td>
<td>1.18</td>
</tr>
</tbody>
</table>

Table A.1

Answers to Calculations:

1) The wt. of the apparatus full of O2 minus the wt. of the apparatus = the apparent wt. of O2.

2) The volume of the bag times the wt./liter of air at the appropriate temperature and pressure = the wt. of air displaced.

Calculations:

1) What is the apparent weight of oxygen that was in the bag?

2) What is the weight of air displaced by the bag?
(3) What is the true weight of the oxygen that was in the bag?

(4) What is the true weight of the carbon dioxide that was in the bag?

(5) What is the ratio by weight of the volume of CO₂(g) to the volume of O₂(g)?

(6) What is the ratio by weight of an individual CO₂(g) particle to an individual O₂(g) particle?

(7) Use a table of atomic weights to calculate the ratio of a mole of CO₂(g) to a mole of O₂(g).

(8) Keeping uncertainty in mind, how do your answers to 5 and 6 compare with the answer to #7?

B. THE VOLUME OF A MOLE OF GAS

One mole of most gases will occupy 22.4 liters if the temperature is 0°C Celsius and the pressure is one atmosphere (760 mm Hg).

C. STANDARD TEMPERATURE AND PRESSURE (S. T. P.)

By arbitrary agreement among scientists throughout the world, standard temperature is 0°C and standard pressure is one atmosphere (760 mm Hg).
D. TOTAL PRESSURE AND PARTIAL PRESSURE

The figure below shows three bulbs of equal volume. The first bulb contains 0.0050 mole of air. The pressure gauge shows the pressure to be 93 mm Hg. The second bulb contains 0.0011 mole of water vapor. The pressure in the second bulb is 20 mm Hg. The third bulb contains 0.0050 mole of air and 0.0011 mole of water vapor. The pressure in the third bulb is 113 mm Hg. This indicates that the pressure exerted by the mixture of gases is the sum of the pressures of the separate components of the mixture.

\[113 \text{ mm Hg} = 93 \text{ mm Hg} + 20 \text{ mm Hg}\]

Because similar experiments yield the same results, we can state: The total pressure is the sum of the pressures exerted by each of the components of the gas mixture. The pressure exerted by each of the gases in a gas mixture is called the partial pressure of that gas. In other words, the partial pressure is the pressure each gas would exert if it were all by itself in the container.
If the 0.0050 mole of air contains 0.0040 mole of N₂ and 0.0010 mole of O₂, what is the partial pressure of O₂ in the first bulb? What is the partial pressure of O₂ in the third bulb? Don't read further until you have answered these two questions.

D.1 Experiment: A QUANTITATIVE INVESTIGATION OF THE REACTION OF MAGNESIUM WITH HYDROCHLORIC ACID

This experiment gives you the opportunity to determine the volume of hydrogen gas produced when a small amount of magnesium reacts with hydrochloric acid. The hydrogen gas is collected over water at room temperature and pressure. The major purpose of this experiment is to determine the volume of dry hydrogen (that is, H₂(g) without any H₂O(g)) that would be produced by one mole of magnesium.

Obtain a piece of magnesium ribbon and carefully cut off a piece 5.00 cm long. Your teacher will give you the weight of 1.000 meter of Mg ribbon. Since the ribbon is of uniform thickness and width, you can calculate the weight of the short piece of Mg you will use. Record your data promptly (see p.79).

Wad the Mg ribbon so that it can be encased in a cage made of fine copper wire. Leave about 5 cm of copper wire to serve as a handle.

The students should be able to answer these two questions before going on.

An alternative would be to use cotton thread in place of the fine copper wire. Have students push the Mg wound with thread into the tube with a pencil and then invert the tube.
Position a ring stand and utility clamp to hold a 50 ml gas measuring tube which has been fitted with a one-hole rubber stopper. Have a 400 ml beaker about two thirds full of tap water handy.

Now tip the gas measuring tube slightly to one side and pour in 10 ml of the hydrochloric acid which is labeled 6 M HCl. With the tube in the same position, carefully fill the tube with water from your wash bottle. While pouring, rinse any acid that may be on the sides of the tube. This will help make certain the liquid in the top of the tube contains very little acid at the very beginning. Bubbles clinging to the sides of the tube can be dislodged by gently tapping the tube.

Hold the copper coil by the handle and insert the cage with the Mg ribbon about 3 cm down the tube. Bend the remaining copper wire over the edge of the tube and clamp it there by inserting the rubber stopper. The tube should be completely filled so that the stopper displaces some water and some water spurts out the hole.

Cover the hole in the stopper with your finger and invert the tube in the 400 ml beaker. Clamp the tube in place. The hydrochloric acid (HCl) is more dense than water and it will dif-
fuse down and react with the metal.

After the reaction stops, wait for about 5 minutes so that the apparatus will come to room temperature. Tap the tube gently to dislodge any bubbles clinging to the side of the tube.

Now cover the hole in the stopper with your finger and transfer the tube to a large cylinder which is almost full of water at room temperature. The acid solution is dilute enough so that it will not hurt you if you rinse your hand with water after the transfer. Raise or lower the tube until the level of the solution inside the tube matches the water level on the outside of the tube. This permits you to measure the volume of the gases (hydrogen and water vapor) at room pressure. The pressure inside the tube equals the pressure outside the tube (room pressure) when the levels are the same. Why?

Read and record the volume with the appropriate uncertainty.

Then remove the gas measuring tube and clean the apparatus.

Finally, record the room pressure. You may want to repeat the experiment if you have enough time.

Your data table should include the

The pressure (force/unit area) must be the same, otherwise the region of greater pressure will cause a lower water level.
following:
Weight of 1.000 meters of Mg ribbon____
Weight of Mg sample used____
Volume of gases____
Temperature of water (room temperature)____
Barometer reading (room pressure)____
Partial pressure due to water vapor at room
temperature____

The following table will help you determine the
calculations

VAPOR PRESSURE OF WATER AT VARIOUS TEMPERATURES
(To three significant figures)

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Pressure (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>17.5</td>
</tr>
<tr>
<td>21</td>
<td>18.6</td>
</tr>
<tr>
<td>22</td>
<td>19.8</td>
</tr>
<tr>
<td>23</td>
<td>21.0</td>
</tr>
<tr>
<td>24</td>
<td>22.4</td>
</tr>
<tr>
<td>25</td>
<td>23.8</td>
</tr>
<tr>
<td>26</td>
<td>25.2</td>
</tr>
<tr>
<td>27</td>
<td>26.7</td>
</tr>
<tr>
<td>28</td>
<td>28.3</td>
</tr>
<tr>
<td>29</td>
<td>30.0</td>
</tr>
<tr>
<td>30</td>
<td>31.8</td>
</tr>
</tbody>
</table>

(1) Find the weight of the Mg you used from the
grams per meter relationship and the length
of your strip of Mg.

(2) How many moles of Mg. did you use?

(3) Assuming the only two gases involved are
H₂(g) and H₂O(g), determine the part of the

142
pressure due to hydrogen gas.

(4) What volume \((V')\) would the \(H_2(g)\) occupy if it were all by itself and the pressure were \(760\) mm Hg (standard pressure)?

(5) What volume \((V'')\) would the \(H_2(g)\) occupy if the pressure were \(760\) mm Hg and the temperature were \(0^\circ\) Celsius? (standard pressure and temperature).

(6) According to your findings, how many liters of \(H_2(g)\) at S. T. P. would be produced by one mole of magnesium?

Exercises for Home, Desk and Lab (HDL)

(1) One volume of nitrogen gas will combine with three volumes of hydrogen gas to produce two volumes of ammonia gas. Use these data and Avogadro's Hypothesis to determine how many molecules of hydrogen combine with one molecule of nitrogen.

How many molecules of ammonia are produced from one molecule of nitrogen?

(2) There are a number of gas mixtures that have the following combining volume ratios: one vol. A + one vol. B = 2 vol. C

If \(a\) = number of atoms in a molecule of A, 
\(b\) = number of atoms in a molecule of B, 
\(c\) = number of atoms in a molecule of C,

a. Show that if \(a\) is even, \(b\) must

room temperature = \(H_2(g)\) pressure.

(4) Convert the volume of \(H_2(g)\) from the original \(H_2(g)\) pressure (calculation #3) to \(760\) mm Hg. Remember a gas occupies the total volume available, so the volume of \(H_2(g)\) and the volume of \(H_2O\) either separately or together will be the "volume of gases."

\[
\text{volume of } \ H_2(g) \ (\text{calculation #3}) \quad (T) \\
(760 \ \text{mm Hg})
\]

(5) Use \(\frac{(V)}{(P)}\) \(\frac{(T)}{}\) = \(\frac{V''}{(T)}\)

(6) wt. of \(Mg\) used

\[
\text{volume } \ B_2(g) \ \text{at S.T.P.} = \frac{\text{wt. of mole } Mg}{x}
\]

Experimentally \(x\) should equal 22.4 liters to three sig. figs.

\((Mg + 2HCl \rightarrow H_2 + MgCl_2)\)

ANSWERS TO QUESTIONS AND PROBLEMS

(1) One volume of nitrogen gas will combine with three volumes of hydrogen gas to produce two volumes of ammonia gas. Therefore, using Avogadro's Hypothesis, one molecule of nitrogen gas will combine with three molecules of hydrogen gas to produce two molecules of ammonia gas.

(2) The data, using Avogadro's Hypothesis, imply:

\[a + b = 2c\]

therefore:
\[b - 2c = a\]

a. No matter what value \(c\) has, \(2c\) must be an even integer. If \(a\) is an even integer, \(b\) must
also be an even integer. (The difference between two even integers is always even.)

b. If \( a \) is odd, the difference between \( 2c \) (an even number) and \( a \) (an odd number in this case) must be odd. If \( a \) is odd, \( b \) must be odd.

(3) a. Only (4) \( \text{HF}_2 \) is not a consistent molecular formula for the substance fluorine.

NOTE: There is no reason to assume that the substance fluorine is an element rather than a compound.

b. The molecular weights and formulas for fluorine that are consistent with the combining volume data are:

1. \( \text{F}_2, 38 \);
2. \( \text{F}_4, 76 \);
3. \( \text{H}_2\text{F}_2, 40 \);
4. \( \text{H}_2\text{F}_4, 78 \).

(3) One volume of hydrogen gas combines with one volume of fluorine gas to produce two volumes of hydrogen fluoride gas (as measured at the same temperature and pressure).

A variety of other types of evidence suggests that hydrogen is an element and its molecules are diatomic (two atoms per molecule).

a. Which one of the following possible molecular formulas for the substance fluorine is not consistent with the volumes that combine? (Use only the data given here; do not presume the molecular formula of hydrogen fluoride.)

\( \begin{align*}
(1) \text{F}_2 & \quad (3) \text{H}_2\text{F}_2 & \quad (5) \text{H}_2\text{F}_4 \\
(2) \text{F}_4 & \quad (4) \text{HF}_2
\end{align*} \)

b. For each formula in part (a) that is consistent with the combining volume data and the formula \( \text{H}_2 \) for hydrogen, calculate the molecular weight indicated by that formula.

c. For each acceptable formula in part (a), predict the molecular formula for the substance hydrogen fluoride.
(4) Four volumes of hydrogen fluoride gas combine with two volumes of the gas dinitrogen difluoride to form four volumes of gas X.

a. According to Avogadro's Hypothesis, how many molecules of X are produced from one molecule of dinitrogen difluoride?

b. If \( a = \) number of atoms in a molecule of hydrogen fluoride, \( b = \) number of atoms in a molecule of dinitrogen difluoride, \( c = \) number of atoms in a molecule of X, write the relation among \( a \), \( b \) and \( c \) appropriate to the combining volumes given.

c. For each of the following possible values of \( a \) and \( b \), calculate the required value of \( c \).

<table>
<thead>
<tr>
<th>( a )</th>
<th>( b )</th>
<th>( c )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>3</td>
</tr>
</tbody>
</table>

\[ 4a + 2b = 4c \]

If \( a = 1 \) and \( b = 2 \), then \( c = 1 \).

d. No odd value of \( b \) is suggested in question (c). Prove that \( b \) must be an even integer.

(5) Oxygen has a density of 1.429 g/L at S. T. P. and its molecular weight is 32.00.

From these data, calculate the volume oc-
(6) a. \( = 22.4 \text{ L} \)

b. \( 6.023 \times 10^{23} \text{ molecules or Avogadro's Number.} \)

(7) Pressure decreased
Volume increased

(8) Temperature decreased
Volume decreased

(9) \( P_{\text{total}} = \text{sum of the partial pressures.} \)
\( \text{P}_{\text{H}} = P_{\text{T}} - \text{P}_{\text{H}_2\text{O}} \)
\( \text{P}_{\text{H}} = 750 \text{ mm Hg} - .25 \text{ mm Hg} \)
\( \text{P}_{\text{H}} = 725 \text{ mm Hg} \)

(10) \( \text{P}_{\text{H}_2\text{O}(g)} \text{ at } 23^\circ \text{C is} \)
\( 23 \text{ mm Hg} \)
\( \text{P}_{\text{O}} = P_{\text{total}} - \text{P}_{\text{H}_2\text{O}(g)} \)
\( \text{P}_{\text{O}} = 800 \text{ mm Hg} - .21 \text{ mm Hg} \)
\( \text{P}_{\text{O}} = 779 \text{ mm Hg} \)

cupied by one mole of oxygen at S. T. P.

(6) a. What volume will a molecular weight of any gas occupy at S. T. P.?

b. How many particles of any gas are in a molecular weight at S. T. P.?

(7) By what factor will one mole of a gas change in volume if the pressure is reduced to 740 mm Hg?

(8) By what factor will one mole of a gas change in volume if the temperature is decreased to 260 degrees Kelvin?

(9) Suppose that 200 ml of hydrogen is collected over water at a temperature of 26°C and a pressure of 750 mm Hg. The pressure of water vapor at 26°C is 25 mm Hg. What is the pressure of the hydrogen gas in the dry state at 26°C?

(10) Suppose that 500 ml of oxygen is collected over water at a temperature of 23°C and a pressure of 800 mm Hg. Calculate the pressure of dry oxygen.
Chapter III: A USEFUL FORM OF $P=kDT$

A  VOLUME, TEMPERATURE RELATIONS
B  PRESSURE, VOLUME RELATIONS
C  COMBINED GAS LAWS
<table>
<thead>
<tr>
<th>SECTION</th>
<th>ROUGH TIME ESTIMATES</th>
<th>EXPERIMENTS</th>
<th>DEMONSTRATIONS</th>
<th>TEACHING AIDS</th>
<th>OTHER STUDENT ACTIVITIES</th>
<th>PROBLEMS</th>
<th>OUTSIDE READING</th>
</tr>
</thead>
<tbody>
<tr>
<td>FUL FORM</td>
<td>Temp.,relations</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1, 2, 3</td>
</tr>
<tr>
<td></td>
<td>pressure, volume</td>
<td>2 Days</td>
<td></td>
<td>Film - Gas pressure and molecular collisions CHEMS</td>
<td>4, 5</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>gas</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6, 7, 8, 9, 10</td>
</tr>
</tbody>
</table>
Chapter III: USEFUL FORM OF P=kDT

In Chapter Eleven of Harvard Project Physics, an ideal gas law, P=kDT was constructed. P=kDT can be further modified and hence more easily applied to problems involving volume, temperature and pressure of gases.

The density (D) of a gas is its mass (m) divided by the volume (V) of the gas:

\[ D = \frac{m}{V} \]

where

- D = density
- m = mass
- V = volume of gas

P=kDT can be written as

\[ P = \frac{(k)m}{V} \]

VOLUME, TEMPERATURE RELATIONS

Equation (2) can be rearranged to

\[ V = \frac{km}{P} \]

Because the mass (m) is directly proportional to the number of particles involved (recall Avogadro's Hypothesis), assuming the number of particles to be constant also assumes the mass to be constant. Then V becomes directly proportional to T (the absolute temperature).

\[ V = kT \]

The volume is directly proportional to the absolute temperature (\( V \propto T \)), a relationship which is referred to as Charles' Law.

Although P=kDT is derived in section 11.5 of Harvard Project Physics, the thought process that accompanies the derivation is difficult. If students press for more detail or if you wish a better understanding of the ideal gas law, you might check chapter nine of PSSC or pp. 65-78 of the students guide for Physics - Chemistry, A Two Year Sequence. (The PSSC CHEMS version published by Portland School Dist. #1 of Portland, Oregon.)

If the instructor prefers, gas law laboratory experiments which are widely available may be appropriate here.
The relation between gas volume and temperature can be determined experimentally.

Table A.1 on the next page shows the results of such an experiment. These results were obtained by holding the pressure and the mass of gas constant and varying the volume and temperature.

It is not possible to obtain data at extremely low temperatures because all gases ultimately condense into liquids. However, the experimentally derived curve may be extrapolated to the point which represents zero volume. The temperature at this point is -273°C. This is designated as absolute zero. This is, theoretically, the lowest possible temperature attainable. Any lower temperature would represent a negative volume. Scientists have recorded temperatures as low as 0.0015 degrees above absolute zero.

The Kelvin scale (°K) is a temperature scale based on an absolute zero; it is often called the Absolute scale (°A). Look at the additional lines marked Celsius and Kelvin (absolute) below Figure A.1. How would you convert Celsius temperatures to Kelvin temperatures? You must add 273° to the °C to obtain °K. To change °K to °C you would subtract 273°
T vs. V for one mole of gas
(P constant at one atmosphere)

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Volume</th>
<th>V/1</th>
</tr>
</thead>
<tbody>
<tr>
<td>°C</td>
<td>°K</td>
<td>liters</td>
</tr>
<tr>
<td>-20</td>
<td>253</td>
<td>20.9</td>
</tr>
<tr>
<td>0</td>
<td>273</td>
<td>22.4</td>
</tr>
<tr>
<td>100</td>
<td>373</td>
<td>30.2</td>
</tr>
</tbody>
</table>

Table A.1.

When temperatures are plotted against corresponding volumes a graph such as that in Figure A.1. is obtained.

VOLUME OF G vs TEMPERATURE

If molecules remained in the gaseous phase, the sample would have zero volume at this temperature.

Figure A.1. = Volume of Gas vs. Temperature
Students should be reminded of HP Ex 29.

It is important to note that the mathematical relationship which represents gas behavior is simplified by using the Kelvin temperature scale because it is easier to work with direct proportions. Inspection of Figure A.1. shows that if we use °K for the X axis, the volume will vary by a factor of 1/273 for each degree change in temperature.

B. PRESSURE, VOLUME RELATIONS

If we hold m and T constant, then

\[(5) \quad P = \frac{1}{V}\]

The volume is inversely proportional to the pressure. \((P = \frac{1}{V})\) is often referred to as Boyle's Law.

C. COMBINED GAS LAWS

Now let's assume we have a closed container with a movable piston (Figure A.2.) to which we will supply a definite volume of gas.

![Movable Piston Cylinder Figure A.2.](image)

The initial conditions will be room temperature and atmospheric pressure. Our initial...
conditions can be designated by \( \frac{PV}{T} = mk \). Now, let's move the piston (change the volume) and apply heat (change the temperature), which in each case would affect the pressure. We now have a new set of conditions for our gas sample, and we will designate these new conditions as \( \frac{P'V'}{T'} = m'k' \).

When we examine the variables \( m \) and \( m' \), we observe that in both cases the mass of the gas remains constant. We did not add or subtract molecules, therefore \( m \) should be equal to \( m' \). Since \( k \) depends on the type of gas, and we did not change the gas in the chamber, \( k \) should equal \( k' \). It follows that

\[
mk = m'k'
\]

Since \( mk = m'k' \):

\[
(6) \quad \frac{PV}{T} = \frac{P'V'}{T'}
\]

This combination of both Charles' and Boyle's Laws is called the combined gas law.

Let's use some algebra and rearrange our equation to find \( V' \) (the volume under new conditions). Upon rearrangement, we obtain:

\[
(7) \quad V' = \left( \frac{PV}{P'V'} \right) (V)
\]

From this we can observe that we have two sets of ratios, one dealing with pressure and one dealing with temperature. Let's apply these
ratios and the equation in a sample problem.

Assuming a sample of O$_2$ gas in the piston apparatus occupies 100 ml at 27°C and 750 mm Hg pressure, what volume will it occupy at standard temperature and pressure? Solution:

1. First, itemize the given data as follows: $P = 750$ mm Hg
   $V = 100$ ml
   $T = 27°C$

Remember, it is essential to use the Kelvin temperature scale, so:

$$T_K = 273 + T_C$$

$$T_K = 273 + 27$$

$$T_K = 300$$

where $T_K =$ Temp. in degrees Kelvin

$$T_C =$ Temp. in degrees Celcius

Our problem states that the new conditions of the gas will be at standard temperature and pressure. Remember standard temperature is 0°C or 273°K and standard pressure is 760 mm Hg.

Therefore, itemize the new conditions as follows:

$$P' = 760$$ mm Hg

$$T' = 273°K$$

$$V' = ?$$

2. Use our generalizations about the behavior of gases to determine temper-
ature and pressure ratios: If pressure increases then volume decreases; therefore, our \( \frac{P}{P'} \) ratio must produce a decrease of volume \( (V) \).

The pressure ratio is:

\[
\frac{750 \text{ mm Hg}}{760 \text{ mm Hg}}
\]

If temperature decreases then volume decreases; therefore, our \( \frac{T}{T'} \) ratio must produce a decrease of volume. Thus the temperature ratio is:

\[
\frac{273^\circ \text{K}}{300^\circ \text{K}}
\]

3. To obtain the final volume \( (V') \), multiply the initial volume \( (V) \) by the temperature and pressure fractions.

The result is:

\[
V' = 100 \text{ ml} \frac{(750 \text{ mm Hg})(273^\circ \text{K})}{(760 \text{ mm Hg})(300^\circ \text{K})}
\]

\[
V' = 89.6 \text{ ml}
\]

4. Or we may use the combined gas law formula, \( \frac{P \cdot V}{T} = \frac{P' \cdot V'}{T'} \),

\[
\frac{(750 \text{ mm Hg})(100 \text{ ml})}{(300^\circ \text{K})} = \frac{(760 \text{ mm Hg})(V')}{(273^\circ \text{K})}
\]

Solving for \( V' \):

\[
V' = \frac{(750 \text{ mm Hg})(100 \text{ ml})(273^\circ \text{K})}{(300^\circ \text{K})(760 \text{ mm Hg})}
\]

\[
V' = 89.6 \text{ ml}
\]
a. The volume of hydrogen changes by the factor 1.92.

b. The pressure of hydrogen changes by the same factor 1.92.

c. The answers to (a) and (b) would be the same if we had used nitrogen gas.

293°K

S. temp. = 273°K

Temp ratio = \frac{273°K}{293°K}

200 ml \times \frac{273}{293} = 186 ml

293°C = 293°K

S. temp. = 273°K

Pressure ratio = \frac{750}{760}

400 ml \times \frac{750}{760} = 394 ml

Exercises for Home, Desk and Lab (HDL)

1. a. A thermometer containing hydrogen is at the temperature of a carbon dioxide-acetone bath (−78°C). The volume of hydrogen at standard atmospheric pressure is read. The thermometer is then moved into a bath of boiling water. The pressure on the hydrogen gas volume is still atmospheric pressure. By what factor does the volume of hydrogen change?

b. If the thermometer had a constant volume, by what factor would the pressure of the hydrogen gas change?

c. If we used nitrogen gas instead of hydrogen in the thermometer, how would it affect the answers to (a) and (b)?

2. A student collects 200 ml of dry hydrogen at a temperature of 20°C. What is the volume of dry hydrogen at standard temperature if the pressure doesn't change?

3. A student collects 400 ml of oxygen at a pressure of 750 mm Hg. What is the volume of oxygen at standard pressure if the temperature doesn't change?

4. A 15.0 liter sample of dry air in a cylinder exerts a pressure of 3.00 atmos-
pheres pressure at a temperature of 25°C. Without change in temperature, a piston is moved in the cylinder until the pressure in the cylinder is lowered to 1.00 atmosphere pressure. What is the new volume of gas in the cylinder?

(5) 100 ml of carbon dioxide gas is collected over water at a temperature of 20°C and 800 mm Hg pressure. What is the volume of dry carbon dioxide at S. T. P?

(6) A 300 ml sample of nitrogen at S. T. P. is brought to 27°C and 800 mm Hg. Calculate the new volume.

(7) A sample of gas occupies 275 ml at 52°C and 720 mm Hg. Correct its volume to S. T. P.

(8) A sample of nitrogen is collected over water at 23°C. The vapor pressure of water at 23° is 21 mm Hg. When the pressure on the sample has been equalized against atmospheric pressure, 750 mm Hg, what is the partial pressure of nitrogen? What will be the partial pressure of nitrogen if the volume is reduced by a factor, 720/760?

(9) A cylinder contains oxygen gas and a small amount of liquid water at a temperature

(4) The new volume of gas is 45 liters.

(5) \[P_{CO_2} = P_{total} - P_{H_2O}\]
   \[P_{CO_2} = 800 - 18\]
   \[P_{CO_2} = 782\ mm\ Hg\]
   \[20°C = 293 K\]
   \[100\ ml \times \frac{273 \times 800}{293} = 97.8\ ml\]

(6) 314 ml

(7) 218 ml

(8) The partial pressure of nitrogen is 729 mm Hg. The partial pressure of nitrogen when the volume is reduced is 770 mm Hg.

(9) The final total pressure is 1376 mm Hg.
Because particles in solids are very close together, we should expect solids to be nearly incompressible. Because of the relatively large distance between gas particles, we should expect them to be very compressible. The distance between particles in liquids is relatively more like solids than gases. Therefore, we should expect liquids to be slightly compressible.

(10) Would you expect the compressibility of solids and gases to differ? Would you expect the compressibility of gases and liquids to differ? Why?

of 25°C. (The vapor pressure of water at 25°C is 23.8 mm Hg.) The total pressure is 700.0 mm Hg. A piston is pushed into the cylinder until the volume is halved. What is the final total pressure if the temperature of the gas doesn't change?
Chapter IV: CHEMICAL EQUATIONS

A  WRITING EQUATIONS FOR REACTIONS

B  TYPES OF CHEMICAL REACTIONS
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A. WRITING EQUATIONS FOR REACTIONS

Scientists have a rapid and concise way of representing the relative amounts of reactants and products involved in chemical reactions. This is done by writing chemical equations. Chemical equations include the usual symbols for the elements or compounds involved and numbers representing the number of moles of each chemical.

In order to predict quantitatively the amounts of reactants needed to produce specific amounts of products, a correctly balanced equation is needed. It is obvious one must know the reactants and products if he is going to balance something involving reactants and products.

A good chemist is one who can predict the products when chemicals (reactants) are put together under given conditions. At this point in the course, however, your experience is not extensive enough for you to be able to predict the products of chemical reactions. Until you become proficient in predicting, you will be provided with the names and formulas of both reactants and products. In the meantime, you
should be looking for some definite patterns as we discuss different kinds of reactions.

General Tasks for Writing Equations

The general tasks for writing and balancing equations are few but need to be adhered to faithfully.

Task #1 -- The first step in writing an equation for a chemical reaction is to determine the reactants used and the products formed.

Task #2 -- The second step is to write the correct chemical formula for each reactant and each product. Reactants go on the left of the equation and are separated from the products by an arrow which is read "yields."

\[ A + B \rightarrow C + D \]

Reactant A plus reactant B yields product C plus product D.

Task #3 -- The law of conservation of mass must be satisfied. In accordance with this law, the number of atoms in the system undergoing change must be constant. A new atom cannot appear on the product side, and no atom from the reactant side may disappear. You may feel the urge to change the formulas of reactants and products as you apply task #3. You must resist this urge because in changing them you are admitting you really don't have the correct
formulas. You may change only the numbers that precede the formulas.

Example: Solid copper will react with an aqueous solution of silver nitrate to produce solid silver plus an aqueous solution of copper nitrate. If you know the respective formulas (Cu, AgNO₃, Ag, Cu(NO₃)₂) you have satisfied task #1.

Let’s satisfy task #2:

\[ \text{Cu} + \text{AgNO}_3 \rightarrow \text{Ag} + \text{Cu(NO}_3\text{)}_2 \]

(Eq.1)

We now must satisfy task #3. We must be certain that our representation is consistent with our belief in the conservation of atoms. We have one mole of Cu on the left and one on the right. Good! Let’s check the Ag. We have one mole of Ag on the left and one on the right. Fine! Now let’s check N.

OOPS!!!

One mole on the left but two on the right.
The subscript number after the parenthesis means that everything inside the parentheses is doubled. Hence two moles of nitrogen atoms and six moles of oxygen atoms as well as one mole of copper atoms are represented by Cu(NO₃)₂.

Let’s put \(\frac{1}{2}\) before the Cu(NO₃)₂. Now we have
\[ \text{Cu} + \text{AgNO}_3 \rightarrow \text{Ag} + \frac{3}{4}\text{Cu(NO}_3)_2 \]  
(Eq. 2)

Now the N balances with one mole on each side. A quick check shows three moles of O on each side. But you complain, we have now represented one mole of Cu on the left and only $\frac{1}{2}$ mole on the right. Perhaps the equation will balance if we put "$\frac{1}{2}$" before the Cu on the left.

\[ \frac{1}{2}\text{Cu} + \text{AgNO}_3 \rightarrow \text{Ag} + \frac{1}{2}\text{Cu(NO}_3)_2 \]  
(Eq. 3)

HURRAY!!

Another check shows that we are firm believers in the conservation of mass and that we have completed all the tasks.

It should be noted here that we could have chosen 1 mole of Cu metal as the amount of reactant consumed in this reaction. We have already seen that we need twice as many molecules of AgNO_3 as molecules of Cu.

Therefore the reaction

\[ \text{Cu} + 2\text{AgNO}_3 \rightarrow \text{Cu(NO}_3)_2 + 2\text{Ag} \]  
(Eq. 4)

is also a chemical reaction in which atoms are conserved. It is just as correct as equation (3). To show this we can multiply equation (3) by 2 in order to obtain equation (4)

All coefficients in an equation may be
multiplied or divided by a common factor, resulting in an equally valid equation.

There are two other tasks that will need attention as more advanced chemical reactions are studied. The first of these is that the number of electrons gained must equal the number lost in the reaction and the second is that the overall charge of the reactants must equal the overall charge of the products. These aspects of balancing equations will be considered when the moon is in the seventh house.

B. TYPES OF CHEMICAL REACTIONS

Chemists have found that most chemical reactions tend to group themselves into four general categories or types of reactions. They are as follows:

Type I: Composition (synthesis) reactions are those in which 2 or more substances react to form a more complex substance. Composition reactions have the general form:

\[ A + B \rightarrow AB \]

Examples:

(1) Sodium + Chlorine \( \rightarrow \) Sodium chloride
    \[ 2Na + Cl_2 \rightarrow 2 NaCl \]

(2) Zinc + Oxygen \( \rightarrow \) Zinc oxide
    \[ 2Zn + O_2 \rightarrow 2ZnO \]

Type II: Decomposition reactions are
those in which a complex substance will break down or separate into two or more less complex substances. You will note that decomposition reactions are the reverse of composition (synthesis) reactions. Decomposition reactions have the general form:

$$AB \rightarrow A + B$$

Examples:

1. Sodium $\rightarrow$ Sodium + Oxygen
   
   Chlorate $\rightarrow$ Chloride
   
   $2 \text{NaClO}_3 \rightarrow 2 \text{NaCl} + 3 \text{O}_2$

2. Mercuric $\rightarrow$ Mercury + Oxygen
   
   Oxide
   
   $2 \text{HgO} \rightarrow 2 \text{Hg} + \text{O}_2$

Type III: Single replacement reactions are those in which one substance is replaced in a given compound by another substance. A single replacement reaction has the general form:

$$A + BX \rightarrow AX + B$$

Examples:

1. Zinc + Hydrochloric $\rightarrow$ Zinc chloride
   
   Acid + Hydrogen
   
   $\text{Zn} + 2 \text{HCl} \rightarrow \text{ZnCl}_2 + \text{H}_2$

2. Chlorine + Sodium $\rightarrow$ Sodium chloride
   
   Iodide + Iodine
   
   $\text{Cl}_2 + 2 \text{NaI} \rightarrow 2 \text{NaCl} + \text{I}_2$

Type IV: Double replacement reactions are those in which two compounds interact and exchange their component parts to form two new
compounds. A double replacement reaction has the general form:

\[ AX + BY \rightarrow AY + BX \]

Example:

1. Sodium + Potassium \( \rightarrow \) Sodium chloride nitrate chloride + Potassium nitrate

\[ \text{NaNO}_3 + \text{KCl} \rightarrow \text{NaCl} + \text{KNO}_3 \]

2. Barium + Hydrochloric \( \rightarrow \) Barium hydroxide acid chloride + water

\[ \text{Ba(OH)}_2 + 2 \text{HCl} \rightarrow \text{BaCl}_2 + 2 \text{H}_2\text{O} \]

Exercises for Home, Desk and Lab (HDL)

Write balanced equations for each of the following chemical reactions. The symbol "↓" means the element or compound is a solid precipitate and the symbol "↑" means the element or compound is a gas. A "△" over the "yield" symbol means the addition of heat energy to the system.

1. \( \text{Fe} + \text{O}_2 \rightarrow \text{Fe}_2\text{O}_3 \)  
2. \( \text{N}_2 + \text{H}_2 \rightarrow \text{NH}_3 \)  
3. \( \text{KClO}_3 \rightarrow \text{KCl} + \text{O}_2 \)  
4. \( \text{SiO}_2 + \text{H}_2\text{F}_2 \rightarrow \text{SiF}_4 \uparrow + \text{H}_2\text{O} \)  
5. \( \text{Ag} + \text{HNO}_3 \rightarrow \text{AgNO}_3 + \text{NO}_2 \uparrow + \text{H}_2\text{O} \)  

Balance the equations for each of the following reactions. Begin on the basis of one mole of the substance underscored.

6. \( \text{Na} + \underline{\text{I}_2} \rightarrow \text{NaI} \)  
7. \( \underline{\text{O}_2} \rightarrow \text{O}_2 \uparrow \)  
8. \( \text{AgNO}_3 + \text{NO}_2 \uparrow + 2\text{H}_2\text{O} \)  
9. \( \text{Na} + \underline{\text{I}_2} \rightarrow 2 \text{NaI} \)
(7) \( \text{C}_2\text{H}_2 + \frac{5}{2} \text{O}_2 \rightarrow \text{H}_2\text{O}^+ + 2 \text{CO}_2^+ \)

(8) \( 2 \text{H}_2 + \text{O}_2 \rightarrow 2 \text{H}_2\text{O}^+ \)

(9) \( \text{Ca} + \text{Cl}_2 \rightarrow \text{CaCl}_2 \)

(10) \( \frac{3}{2} \text{NO}_2 + \frac{1}{2} \text{H}_2\text{O} \rightarrow \text{HNO}_3 + \frac{1}{2} \text{NO}^+ \)

You may wish to supply more equations for balancing practice.

(11) \( 2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{NaOH} + \text{H}_2^+ \)

(12) \( 2\text{Ca(OH)}_2 + 2\text{H}_3\text{PO}_4 \rightarrow \text{Ca}_3(\text{PO}_4)_2 + 6\text{H}_2\text{O} \)

(13) \( \text{Ba(NO}_3)_2 + \text{Na}_2\text{SO}_4 \rightarrow \text{BaSO}_4 \downarrow + 2\text{NaNO}_3 \)

(14) \( (\text{NH}_4)_2\text{SO}_4 + 2\text{KOH} \rightarrow \text{K}_2\text{SO}_4 + 2\text{H}_2\text{O} + 2\text{NH}_3 \uparrow \)

(15) \( \text{Cl}_2 + \text{MgBr}_2 \rightarrow \text{MgCl}_2 + \text{Br}_2 \)

(16) \( \text{C}_3\text{H}_8 + 5\text{SO}_2 \rightarrow 3\text{CO}_2^+ + 4\text{H}_2\text{O} \)

Write a balanced equation and indicate the reaction type (composition, decomposition, single or double replacement) for each of the following word equations:

(You may wish to consult the Handbook of Physics and Chemistry in order to obtain the correct formulas for the reactants and products involved in these reactions.)

(11) Sodium + Water \rightarrow Sodium + Hydrogen hydroxide

(12) Calcium hydroxide + Phosphoric acid \rightarrow Calcium phosphate + Water

(13) Barium nitrate + Sodium sulfate \rightarrow Barium sulfate + Sodium nitrate

(14) Ammonium sulfate + Potassium hydroxide \rightarrow Potassium sulfate + water + Ammonia gas

(15) Chlorine + Magnesium bromide \rightarrow Magnesium chloride + Bromine

Propane, a commercially bottled gas, has the formula \( \text{C}_3\text{H}_8 \). When it is burned completely the reaction products are carbon dioxide and water vapor.

(16) Write the balanced equation for the
combustion of propane.

(17) One mole of propane vapor produces how many moles of water vapor when burned?

(18) How many moles of oxygen would be needed in order to burn 0.2 moles of propane?

(19) How many moles of propane would be needed to produce one mole of carbon dioxide?

(20) How many molecules of oxygen would be needed in order to burn one mole of propane?

(17) 4 moles of water vapor

(18) 1 mole of oxygen

(19) \( \frac{1}{3} \) mole of propane

(20) \( 3.01 \times 10^{54} \) molecules of oxygen
Chapter V: ELECTRICAL NATURE OF MATTER

A  PURE SUBSTANCES
   Read sections 5-1 through 5-1.3 CHEMS

B  SOLUTIONS
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5-4.2 Precipitation reaction in aqueous solutions

5-4.3 Electrical conductivity of solids

5-4.4 Ionic solids
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E  HYDROGEN - A FAMILY BY ITSELF
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Chapter VII: ENERGY EFFECTS IN CHEMICAL REACTIONS

A
HEAT AND CHEMICAL REACTIONS
Read sections 7-1 through 7-1.4
CHEMS

A.1 Experiment: A STUDY OF REACTIONS
(CEx 12)

A.2 Experiment: THE HEAT OF REACTION
(CEx 13)

B
THE LAW OF CONSERVATION OF ENERGY
Read sections 7-2 through 7-2.4
CHEMS

C
THE ENERGY STORED IN A MOLECULE
Read sections 7-3 through 7-3.2
CHEMS

D
THE ENERGY STORED IN A NUCLEUS
Read sections 7-4 through 7-4.1
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<td>3 days</td>
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<td>Scientific American Sept. 1954</td>
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<td>Ex 13 The heat of reaction</td>
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<td>Scientific American July 1957 pp. 72-88</td>
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<td>Scientific American Sept. 1956 pp. 82-91</td>
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Chapter VIII: RATES OF REACTIONS

A  FACTORS AFFECTING REACTION RATES
   Read sections 8-1 through 8-1.5
   CHEMS

A.1  Experiment: A STUDY OF REACTION RATES
     (CEx 14)

B  THE ROLE OF ENERGY IN REACTION RATES
   Read sections 8-2 through 8-2.4
   CHEMS
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<td>Scientific American Nov. 1957 pp. 117-126</td>
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Ex. 1: 1-8 incl.  
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Chapter IX: EQUILIBRIUM

A  QUALITATIVE ASPECTS OF EQUILIBRIUM
Read sections 9-1 through 9-1.7
CHEMS

A.1 Experiment: CHEMICAL EQUILIBRIUM
(CEx 15)

B  QUANTITATIVE ASPECTS OF EQUILIBRIUM
Read sections 9-2 through 9-2.4
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<td>Ex 15 Chemical equilibrium</td>
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<td>Ex. 2 7-16 incl.</td>
<td>Equilibrium by M.J. Sienko, Benjamin, Inc. 1964</td>
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Chapter X: SOLUBILITY

A  SOLUBILITY: A CASE OF EQUILIBRIUM
Read sections 10-1 through 10-1.4
CHEMS

B  AQUEOUS SOLUTIONS
Read sections 10-2 through 10-2.6
CHEMS

B.1 Experiment: DETERMINATION OF THE
SOLUBILITY PRODUCT CONSTANT OF
SILVER ACETATE
(CEx 16)
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Chapter XI: ACID - BASE

A  ELECTROLYTES - STRONG OR WEAK
Read sections 11-1 through 11-1.3
CHEMS

A.1 Experiment: THE HEAT OF SOME
ACID-BASE REACTIONS
(CEx 17)

B  EXPERIMENTAL INTRODUCTION TO
ACIDS AND BASES
Read sections 11-2 through 11-2.7
CHEMS

B.1 Experiment: DETERMINATION OF THE
HYDROGEN ION CONCENTRATION OF
SOLUTIONS USING INDICATORS
(CEx 18)

C  STRENGTHS OF ACIDS
Read sections 11-3 through 11-3.5
CHEMS

C.1 Experiment: REVERSIBLE CHEMICAL
REACTIONS
(CEx 19)
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Chapter XII: OXIDATION - REDUCTION

A ELECTROCHEMICAL CELLS
Read sections 12-1 through 12-1.4
CHEMS

A.1 Experiment: INTRODUCTION TO OXIDATION REACTIONS
(CEx 20)

A.2 Experiment: ELECTROCHEMICAL CELLS
(CEx 21)

B ELECTRON TRANSFER AND PREDICTING REACTIONS
Read sections 12-2 through 12-2.5
CHEMS

C BALANCING OXIDATION-REDUCTION REACTIONS
Read sections 12-3 through 12-3.3
CHEMS

D ELECTROLYSIS
Read section 12-4
CHEMS
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Chapter XIII: STOICHIOMETRY

A PATTERN FOR STOICHIOMETRIC CALCULATIONS
Read section 13-1
CHEMS

THE MANUFACTURE OF SULFURIC ACID
Read sections 13-2 through 13-2.5
CHEMS

B.1 Experiment: REACTIONS BETWEEN IONS
(CEx 22, Parts I and II)

B.2 Experiment: QUANTITATIVE TITRATION
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