To Cindy
Preface to the Second Edition

It is heartening to hear of your book being read and enjoyed, literally cover to cover, by individuals ranging from talented high-school science students to Professors Emeritus of the English language. Even better to hear that you have a chance to improve that book, based upon the above comments, comments by reviewers, and the experience gained from working with the text.

In this edition of *The Organic Chem Lab Survival Manual*, the section on notebooks and handbooks have been expanded to include typical notebook pages and actual handbook entries along with interpretation. There are new notes on cleaning and drying glassware, and how to find a good recrystallization solvent. Once their samples are purified, students may now find directions for taking a melting point with the Thomas–Hoover apparatus. Washing has been given the same importance as extraction, and a few more trouble spots—taking the pH of an organic layer, for one—have been smoothed. There are additional instructions on steam distillation using external sources of steam. Simple manometers, coping with air leaks, and the correct use of a pressure–temperature nomograph enhance the section on vacuum distillation. Refractometry has been added, as well as—by special request—sections on the theory of extraction and distillation, including azeotropes and azeotropic distillation, and, I believe, the first application of the Clausius–Clapyron equation as a bridge for getting from Raoult’s Law (pressure and mole fraction) to the phase diagram (temperature and mole fraction).

Many people deserve credit for their assistance in producing this edition: my students, for helping me uncover what was lacking in the previous edition, with Mr. Ronald Pohadsky and Mr. Barry Eggleston making specific suggestions while working in the laboratory. A special thanks to Professor G.J. Janz, director of the Molten Salts Data Center at the Rensselaer Polytechnic Institute for his review of the physical chemistry sections of this edition, and to Professors Henry Hollinger and A. Rauf Imam for their help during the initial phases of that work. I would also like to thank

William Epstein  
University of Utah  
Rudolph Goetz  
Michigan State University
Clelia W. Mallory
University of Pennsylvania
J. Wolinsky
Purdue University

for their valuable comments and suggestions in making this edition more useful for students of organic chemistry laboratory.

Finally, I'd like to thank Mr. Dennis Sawicki, Chemistry Editor at John Wiley & Sons, first, for one of the nicest birthday presents I've gotten in a while, and second, for his encouragement, guidance, and patience at some troubling points in the preparation of this edition. Ms. Dawn Reitz, Production Supervisor, Ms. Ann Meader, Supervising Copy Editor, and Mr. Glenn Petry, Copy Editor deserve a great deal of credit in bringing this second edition about.

J. W. Zubrick
Hudson Valley Community College
April 3, 1987
Preface to the First Edition

Describe, for the tenth time, an instrument not covered in the laboratory book, and you write a procedure. Explain, again and again, operations that are in the book, and you get a set of notes. When these produce questions you revise until the students, not you, finally have it right. It you believe that writing is solidified speech — with the same pauses, the same cadences — then a style is set. And if you can still laugh, you write this book.

This book presents the basic techniques in the organic chemistry laboratory with the emphasis of doing the work correctly the first time. To this end, examples of what can go wrong are presented with admonishments, often bordering on the outrageous, to forestall the most common of errors. This is done in the belief that it is much more difficult to get into impossible experimental troubles once the student has been warned of the merely improbable ones. Complicated operations, such as distillation and extraction, are dealt with in a straightforward fashion, both in the explanations and in the sequential procedures.

The same can be said for the sections concerning the instrumental techniques of GC, IR, NMR, and HPLC. The chromatographic techniques of GC and HPLC are presented as they relate to thin-layer and column chromatography. The spectroscopic techniques depend less on laboratory manipulation and so are presented in terms of similarities to the electronic instrumentation of GC and HPLC techniques (dual detectors, UV detection in HPLC, etc.). For all techniques, the emphasis is on correct sample preparation and correct instrument operation.

Many people deserve credit for their assistance in producing this textbook. It has been more than a few years since this book was first written, and a list of acknowledgements would approach the size of a small telephone directory — there are too many good people to thank directly.

For those who encouraged, helped, and constructively criticized, thanks for making a better book that students enjoy reading and learning from.

I'd like to thank the hundreds of students who put up with my ravings, rantings, put-ons, and put-downs, and thus taught me what it was they needed to know, to survive organic chemistry laboratory.

A special thanks to Dr. C.W. Schimelpfenig, for encouragement over many years when there was none, and whose comments grace these pages; Dr. D.L. Carson, whose comments also appear, for his useful criticism concerning the
presentation; Drs. R.A. Bailey, S.C. Bunce, and H.B. Hollinger for their constant support and suggestions; Dr. Mark B. Freilich, whose viewpoint as an inorganic chemist proved valuable during the review of manuscript; and Dr. Sam Johnson, who helped enormously with the early stages of the text processing. I also thank Christopher J. Kemper and Keith Miller for their valuable comments on the instrumental sections of the book.

Finally, I'd like to thank Clifford W. Mills, my patron saint at John Wiley & Sons, without whose help none of this would be possible, and Andrew E. Ford, Jr., vice president, for a very interesting start along this tortured path to publication.
It is common to find instructors railing against poor usage and complaining that their students cannot do as much as to write one clear, uncomplicated, communicative English sentence. Rightly so. Yet I am astonished that the same people feel comfortable with the long and awkward passive voice, the pompous “we” and the clumsy “one,” and that damnable “the student,” to whom exercises are left as proofs. These constructions, which appear in virtually all scientific texts, do not produce clear, uncomplicated, communicative English sentences. And students do learn to write, in part, by following example.

I do not go out of my way to boldly split infinitives, nor do I actively seek prepositions to end sentences with. Yet by these constructions alone, I may be viewed by some as aiding the decline in students’ ability to communicate.

E.B. White, in the second edition of The Elements of Style (Macmillan, New York, 1972, p. 70), writes

Years ago, students were warned not to end a sentence with a preposition; time, of course, has softened that rigid decree. Not only is the preposition acceptable at the end, sometimes it is more effective in that spot than anywhere else. “A claw hammer, not an axe, was the tool he murdered her with.” This is preferable to “A claw hammer, not an ax, was the tool with which he murdered her.” Some infinitives seem to improve on being split, just as a stick of round stovewood does. “I cannot bring myself to really like the fellow.” The sentence is relaxed, the meaning is clear, the violation is harmless and scarcely perceptible. Put the other way, the sentence becomes stiff, needlessly formal. A matter of ear.

We should all write as poorly as White.

With the aid of William Strunk and E.B. White in The Elements of Style and that of William Zinsser in On Writing Well, Rudolph Flesch in The ABC of Style, and D.L. Carson, whose comments appear in this book, I have tried to follow some principles of technical communication lately ignored in scientific texts: use the first person, put yourself in the reader’s place, and, the best for last, use the active voice and a personal subject.
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Seldom does one have the opportunity to read and use a textbook that is completely useful, one that does not need substitutions and deletions. Zubrick's book is this type of resource for undergraduate organic students and their laboratory instructors and professors. I must heartily recommend this book to any student taking the first laboratory course in organic chemistry.

The Organic Chem Lab Survival Manual is filled with explanations of necessary techniques in much the same way that advanced techniques have been presented in books by Wiberg, Lowenthal, Newman, and Gordon and Ford. In larger universities, The Survival Manual is a valuable supplement to most laboratory manuals. It provides explanations that many graduate teaching assistants do not take time to give to their classes. Most teaching assistants of my acquaintance appreciate Zubrick's book because it supports their discussions during recitations (when each student has a personal copy), and it refreshes their memories of good techniques they learned and must pass on to a new generation of undergraduates.

The book is addressed to the undergraduate student audience. The informal tone appeals to most laboratory students. The illustrations are delightful. The use of different type fonts is effective for emphasis. Also, Zubrick always explains why the particular sequence of operations is necessary, as well as how to manipulate and support the apparatus and substances. This is a definite strength.

This book is an evolutionary product: Over the span of a decade, professors at major universities and liberal arts colleges have made suggestions for minor changes and improvements. I count myself fortunate to have used the forerunners, which have been published since 1973.

A large quantity of useful information has been collected, well organized, and presented with great care. This book is the handiwork of a master teacher.

C.W. Schimelpfenig
Dallas Baptist University
Dallas, Texas

The Organic Chem Lab Survival Manual is a book I have known about for a number of years in a variety of developmental stages. As it progressed, I watched with interest as Jim Zubrick struggled to achieve a balance between merely conveying information — what most books do — and conveying that...
information efficiently to its very human audience. On the one hand, Jim insisted that his book contain all the necessary scientific detail; on the other hand, he also insisted that a "how to" book for organic chemistry lab need not be written in the dull and confusing prose which so often passes as the *lingua franca* of science. This book demonstrates that he has achieved both goals in admirable fashion.

In fact, *The Survival Manual* succeeds very well in following Wittgenstein's dictum that "everything that can be thought at all, can be thought clearly. Anything that can be said can be said clearly." It also follows the advice of Samuel Taylor Coleridge to avoid pedantry by using only words "suitable to the time, place, and company."

Although some few readers may take umbrage with this book because it is not, atypically couched in the language of a typical journal article, similar people no doubt also complained when William Strunk published *Elements of Style* in 1919. For Strunk also broke with tradition. Most other writing texts of the day were written in the convoluted language of the nineteenth century, and the material they contained consisted largely of lists of arcane practices, taboos, and shibboleths—all designed to turn students into eighteenth-century writers.

From Strunk's point of view, such texts were less than desirable for several major reasons. First, the medicine they offered students had little to do with the communication process itself; second, it had little to do with current practice; and third, taking the medicine was so difficult that the cure created more distress in the patients than did the disease itself.

Jim Zubrick proves in this book that he understands, as did Strunk, that learning reaches its greatest efficiency in situations where only that information is presented which is directly related to completing a specific task. In an environment fraught with hazards, efficiency of this sort becomes even more necessary.

*The Survival Manual* is an excellent book because it speaks to its audience's needs. Always direct—if sometimes slightly irreverent—the book says clearly what many other books only manage to say with reverent indirection. It never forgets that time is short or that the learning curve rises very slowly at first. The prose is straightforward, easy to understand, and is well supported by plentiful illustrations keyed to the text. It is also technically accurate and technically complete, but it always explains matters of laboratory technology in a way designed to make them easily understandable to students in a functional context.

All of these characteristics related to communication efficiency will natu-
rally make the laboratories in which the book is used safer labs; the improved understanding they provide serves as natural enhancement to the book’s emphatic and detailed approach to laboratory safety.

Most important, however, all the elements of *The Survival Manual* come together in focusing on the importance of task accomplishment in a way which demonstrates the author’s awareness that communication which does not efficiently meet the needs of its audience is little more than *pedantry* unsuitable to the time, place, and company.

David L. Carson
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Safety First, Last and Always
The organic chemistry laboratory is potentially one of the most dangerous of undergraduate laboratories. That is why you must have a set of safety guidelines. It is a very good idea to pay close attention to these rules, for one very good reason:

The penalties are only too real.

Disobeying safety rules is not at all like flouting many other rules. You can get seriously hurt. No appeal. No bargaining for another 12 points so you can get into medical school. Perhaps as a patient, but certainly not as a student. So, go ahead. Ignore these guidelines. But remember —

You have been warned!

1. **Wear your goggles.** Eye injuries are extremely serious and can be mitigated or eliminated if you keep your goggles on at all times. And I mean over your eyes, not on top of your head or around your neck. There are several types of eye protection available, some of it acceptable, some not, according to local, state and federal laws. I like the clear plastic goggles that leave an unbroken red line on your face when you remove them. Sure, they fog up a bit, but the protection is superb. Also, think about getting chemicals or chemical fumes trapped under your contact lenses before you wear them to lab. Then don’t wear them to lab. Ever.

2. **Touch not thyself.** Not a Biblical injunction, but a bit of advice. You may have just gotten chemicals on your hands, in a concentration that is not noticeable, and sure enough, up go the goggles for an eye wipe with the fingers. Enough said.

3. **There is no “away.”** Getting rid of chemicals is a very big problem. You throw them from here, and they wind up poisoning someone else. Now there are some laws to stop that from happening. The rules were really designed for industrial waste, where there are hundreds of gallons of waste that have the same composition. In a semester of organic lab there will be much smaller amounts of different materials. Waste containers could be provided for everything, but this is not practical. If you don’t see the waste can you need, ask your instructor. When in doubt, ask.

4. **Bring a friend.** You must never work alone. If you have a serious
accident and you are all by yourself, you might not be able to get help before you die. Don’t work alone, and don’t work at unauthorized times.

5. **Don’t fool around.** Chemistry is serious business. Don’t be careless or clown around in lab. You can hurt yourself or other people. You don’t have to be somber about it; just serious.

6. **Drive defensively.** Work in the lab as if someone else were going to have an accident that might affect you. Keep the goggles on because *someone else* is going to point a loaded, boiling test tube at you. *Someone else* is going to spill hot, concentrated acid on your body. Get the idea?

7. **Eating, drinking, or smoking in lab.** Are you kidding? Eat in a chem lab? Drink in a chem lab?? Smoke, and blow yourself up????

8. **Keep it clean.** Work neatly. You don’t have to make a fetish out of it, but try to be neat. Clean up spills. Turn off burners or water or electrical equipment when you’re through with them.

9. **Where it’s at.** Learn the location and proper use of the fire extinguishers, fire blankets, safety showers, and eyewashes.

10. **Making the best-dressed list.** No open-toed shoes, sandals, or canvas-covered footwear. No loose-fitting cuffs on the pants or the shirts. Nor are dresses appropriate for lab, guys. Keep the mid-section covered. Tie back that long hair. And, a small investment in a lab coat can pay off, projecting that extra professional touch. It gives a lot of protection too. Consider wearing disposable gloves. Clear polyethylene ones are inexpensive, but the smooth plastic is slippery, and there’s a tendency for the seams to open when you least expect it. Latex examination gloves keep the grip and don’t have seams, but they cost more. Gloves are not perfect protectors. Reagents like bromine can get through and cause severe burns. They’ll buy you some time though, and can help mitigate or prevent severe burns.

11. **Hot under the collar.** Many times you’ll be asked or told to heat something. Don’t just automatically go for the Bunsen burner. That way lies fire. Usually —

    **No Flames!**

Try a hot plate, try a heating mantle (see Chapter 13, “Sources of Heat”). But try to stay away from flames. Most of the fires I’ve had to put out started when some bozo decided to heat some flammable sol-
vent in an open beaker. Sure, there are times when you’ll HAVE to use a flame, but use it away from all flammables and in a hood (Fig. 1), and only with the permission of your instructor.

12. **Work in the Hood.** A hood is a specially constructed workplace that has, at the least, a powered vent to suck noxious fumes outside. There’s also a safety glass or plastic panel you can slide down as protection from exploding apparatus (Fig. 1). If it is at all possible, treat every chemical (even solids) as if toxic or bad smelling fumes came from it, and carry out as many of the operations in the organic lab as you can inside a hood, unless told otherwise.

13. **Keep your fingers to yourself.** Ever practiced “finger chemistry?” You’re unprepared so you have a lab book out, and your finger points to the start of a sentence. You move your finger to the end of the first line, and do that operation—

```
"Add this solution to the beaker containing the ice-water mixture"
```

And WOOSH! Clouds of smoke. What happened? The next line reads—

![Fig. 1 A typical hood.](image-url)
“very carefully as the reaction is highly exothermic.”

But you didn’t read that line, or the next, or the next. So you are a danger to yourself and everyone else. Read and take notes on any experiment before you come to lab (see Chapter 2, “Keeping a Notebook”).

14. **What you don’t know can hurt you.** If you are not sure about any operation, or you have any question about handling anything, *please* ask your instructor before you go on. Get rid of the notion that asking questions will make you look foolish. Following this safety rule may be the most difficult of all. Grow up. Be responsible for yourself and your own education.

15. **Blue Cross or Blue Shield?** Find out how you would get medical help, if you needed it. Sometimes during a summer session, the school infirmary is closed and you would have to be transported to the nearest hospital.

These are a few of the safety guidelines for an organic chemistry laboratory. You may have others particular to your own situation.

**ACCIDENTS WILL NOT HAPPEN**

That’s an attitude you might hold while working in the laboratory. You are NOT going to do anything, or get anything done to you, that will require medical attention. If you do get cut, and the cut is not serious, wash the area with water. If there’s serious bleeding, apply direct pressure with a clean, preferably sterile, dressing. For a minor burn, let cold water run over the burned area. For chemical burns to the eyes or skin, flush the area with lots of water. In every case, get to a physician if at all possible.

If you have an accident, *tell your instructor immediately. Get help!* This is no time to worry about your grade in lab. If you put grades ahead of your personal safety, be sure to see a psychiatrist after the internist finishes.
Keeping a Notebook
A research notebook is perhaps one of the most valuable pieces of equipment you can own. With it you can duplicate your work, find out what happened at leisure, and even figure out where you blew it. General guidelines for a notebook are:

1. The notebook must be bound permanently. No loose leaf or even spiral-bound notebooks will do. It should have a sewn binding so that the only way pages can come out is to cut them out. (8 1/2 x 11 in. is preferred).

2. Use waterproof ink! Never pencil! Pencil will disappear with time, and so will your grade. Cheap ink will wash away and carry your grades down the drain. Never erase! Just draw one line through your errors so that they can still be seen. And never, never, never cut any pages out of the notebook!

3. Leave a few pages at the front for a table of contents.

4. Your notebook is your friend, your confidant. Tell it:
   a. What you have done. Not what it says to do in the lab book. What you, yourself, have done.
   b. Any and all observations: color changes, temperature rises, explosions . . . , anything that occurs. Any reasonable explanation why whatever happened, happened.

5. Skipping pages is extremely poor taste. It is NOT done!

6. List the IMPORTANT chemicals you’ll use during each reaction. You should include USEFUL physical properties: the name of the compound, molecular formula, molecular weight, melting point, boiling point, density, and so. The CRC Handbook of Chemistry and Physics, originally published by the Chemical Rubber Company and better known as the CRC Handbook, is one place to get this stuff (see Chapter 3, “Interpreting a Handbook”).

Note the qualifier “USEFUL.” If you can’t use any of the information given, do without it! You look things up before the lab so you can tell what’s staring back out of the flask at you during the course of the reaction.
Your laboratory experiments can be classified to two major types: a technique experiment or a synthesis experiment. Each requires different handling.

**A TECHNIQUE EXPERIMENT**

In a technique experiment, you get to practice a certain operation before you have to do it in the course of a synthesis. Distilling a mixture of two liquids to separate them is a typical technique experiment.

Read the following handwritten notebook pages with some care and attention to the typeset notes in the margin. A thousand words are worth a picture or so (Figs. 2–4).

**Notebook Notes**

1. Use a descriptive title for your experiment. *Distillation*. This implies you’ve done all there is in the entire field of distillation. You haven’t? Perhaps all you’ve done is *The Separation of a Liquid Mixture by Distillation*. Hmmmmmm.

2. Writing that first sentence can be difficult. Try stating the obvious.

3. There are no large blank areas in your notebook. Draw sloping lines through them. Going back to enter observations after the experiment is over is not professional. Initial and date pages anytime you write anything in your notebook.

4. Note the appropriate changes in verb tense. Before you do the work, you might use the present or future tense writing about something that hasn’t happened yet. During the lab, since you are supposed to write what you’ve actually done just after the time you’ve actually done it, a simple past tense is sufficient.

**A SYNTHESIS EXPERIMENT**

In a synthesis experiment, the point of the exercise is to prepare a clean sample of the product you want. All the operations in the lab (e.g., distillation,
The Separation of a Liquid Mixture by Distillation

Distillation is one of the methods of separation and purification of liquids. We will be given an unknown liquid mixture and will have to separate it by distillation.

After we get the unknown, should deposit wash dry the liquid over anhydrous magnesium sulfate. The setup is as detailed in the laboratory manual with some changes:

![Diagram of distillation apparatus]

We will be using Thunwell lenses and will not need Varilux. Vacuum adapter clamped at angle, rotated toward me in order to make it easy to change flask.

Fig. 2 Notebook entry for a technique experiment (1).
Separation of a Liquid Mixture (cont'd)

Obtained liquid unknown #20, from instructor & placed it over a slight XS of anhydrous magnesium sulfate. Set up distillation apparatus as described (p.6)

Started with the smallest flask to collect fore-run as suggested by instructor. Filtered unknown into distilling flask with long stem funnel. Set heat controller to 50. Mixture beginning to boil.

Liquid condensed on thermometer & temperature reading shot up to 79°C and stabilized at 81°C in a few seconds. Collected a 2 ml as fore-run. Will discard this later. Dropped Thermowell to remove heat to stop distillation and change receiving flashes. Started heating again.

Collected liquid boiling from 81 to 83°C. Charged receive as above. When new material came over, Thermometer read 92°C(!) for a few minutes (al) then distillation stopped. Temperature began dropping! Turned heat up (70). Mixture started to boil again and liquid came over @ 123°C. Collected a little of this then changed receive as above. Shock distilling flask a little & added boiling stone before heating. Had to label flashes. So many of these.

Fig. 3 Notebook entry for a technique experiment (2).
### Separation of a Liquid Mixture (cont'd)

<table>
<thead>
<tr>
<th>Flask</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>free run</td>
</tr>
<tr>
<td>2</td>
<td>81°C - 83°C fraction</td>
</tr>
<tr>
<td>3</td>
<td>82°C - 123°C change-over</td>
</tr>
<tr>
<td>4</td>
<td>120°C - 123°C fraction</td>
</tr>
<tr>
<td>5</td>
<td>&gt;123°C distilling flask residue</td>
</tr>
</tbody>
</table>

Small amount of liquid left over in the boiling flask can't get over. Dangerous to heat to dryness. Stopped distillation after collecting fraction from 120-123°C (Flask #4).

Cooled distilling flask and poured contents into a 50 ml erlenmeyer (Flask #5).

Checked cork stoppers for security. Have permission to store flasks properly labelled, in book until next lab.

**Fig. 4** Notebook entry for a technique experiment (3).
recrystallization, etc.) are just means to this end. The preparation of 1-bromobutane is a classic synthesis, and is the basis of the next series of handwritten notebook pages.

Pay careful attention to the typeset notes in the margins, as well as the handwritten material. Just for fun, go back and see how much was written for the distillation experiment, and how that is handled in this synthesis (Figs. 5–10).

Once again, if your own instructor wants anything different, do it. The art of notebook keeping has many schools — follow the perspective of your own.

**Notebook Notes**


2. Do you see a section for unimportant side reactions? No. Then don’t include any.

3. In this experiment, we use a 10% aqueous sodium hydroxide solution as a wash (see Chapter 11, “Extraction and Washing”), and anhydrous calcium chloride as a drying agent (see Chapter 7, “Drying Agents”). These are not listed in the Table of Physical Constants. They are neither reactants nor products. Every year, however, somebody always lists the physical properties of solid sodium hydroxide, calcium chloride drying agent, and a bunch of other reagents that have nothing to do with the main synthetic reaction. I’m specially puzzled by the listing of solid sodium hydroxide in place of the 10% solution.

4. **Theoretical yield** (not yeild) calculations always seem to be beyond the ken of a lot of you, even though these are exercises right out of the freshman year chemistry course. Yes, we do expect you to remember some things from courses past; the least of which is where to look this up. I’ve put a sample calculation in the notebook (Fig. 7), that gets the mass (g) of the desired product (1-bromobutane) from the volume (ml) of one reactant (1-butanol). Why from the 1-butanol and not from the sulfuric acid or sodium bromide? It’s the 1-butanol we are trying to convert to the bromide, and we use a molar excess (often abbreviated XS) of everything else. The 1-butanol is, then, the **limiting reagent**; the reagent
Synthesis of 1-Bromobutane

We will be preparing 1-bromobutane as follows:

(1) \( \text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} + \text{NaBr} \xrightarrow{\text{H}_2\text{SO}_4} \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{Br} + \text{H}_2\text{O} + \text{NaOH} \)

Side reactions:

(2) \( \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} \xrightarrow{\text{H}_2\text{SO}_4} (\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2)_2\text{O} \)
(3) \( \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} \xrightarrow{\text{H}_2\text{SO}_4} \text{CH}_3\text{CH}_2\text{CH} = \text{CH}_2 + \text{H}_2\text{O} \)
(4) \( \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} \xrightarrow{\text{H}_2\text{SO}_4} \text{CH}_3\text{CH} = \text{CH} - \text{CH}_3 + \text{H}_2\text{O} \)

Table of Physical Constants:

<table>
<thead>
<tr>
<th>Name</th>
<th>Formula</th>
<th>M.W.</th>
<th>Dens.</th>
<th>M.R.</th>
<th>B.P.</th>
<th>Water</th>
<th>Ether</th>
<th>Conc. HSO4</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Butanol</td>
<td>( \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} )</td>
<td>74.12</td>
<td>0.9098</td>
<td>-39.5</td>
<td>117.5</td>
<td>s</td>
<td>s</td>
<td>u</td>
<td>u</td>
</tr>
<tr>
<td>Sulphuric Acid</td>
<td>( \text{H}_2\text{SO}_4 )</td>
<td>98.08</td>
<td>1.8111</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium Bromide</td>
<td>( \text{NaBr} )</td>
<td>102.9</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Bromobutane</td>
<td>( \text{CH}_3\text{CH}_2\text{CH}_2\text{Br} )</td>
<td>137.03</td>
<td>1.2764</td>
<td>-112.3</td>
<td>107.3</td>
<td>i</td>
<td>s</td>
<td>i</td>
<td>s</td>
</tr>
<tr>
<td>p-Dibromobutane</td>
<td>( \text{(CH}_3\text{CH}_2\text{CH}_2\text{Br})_2 )</td>
<td>130.23</td>
<td></td>
<td>142</td>
<td>2.5</td>
<td>s.s.</td>
<td>s</td>
<td>i</td>
<td>s</td>
</tr>
<tr>
<td>trans-2-Butenone</td>
<td>( \text{CH}_3\text{CH} = \text{CHCH}_2\text{CH}_3 )</td>
<td>56.10</td>
<td></td>
<td></td>
<td>1</td>
<td>i</td>
<td>i</td>
<td>i</td>
<td>i</td>
</tr>
<tr>
<td>cis-2-Butenone</td>
<td>( \text{CH}_3\text{CH} = \text{CHCH}_2\text{CH}_3 )</td>
<td>56.10</td>
<td></td>
<td></td>
<td>-5</td>
<td>i</td>
<td>v.s.</td>
<td>i</td>
<td>i</td>
</tr>
<tr>
<td>1-Butene</td>
<td>( \text{CH}_3\text{CH} = \text{CHCH}_2\text{CH}_3 )</td>
<td>56.10</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5 Notebook entry for a synthesis experiment (1).
Calculations from freshman chemistry.

**Synthesis of 1-Bromobutane (cont'd.)**

Theoretical yield of 1-bromobutane based on 17 mL 1-butanol.

\[
17 \text{ mL } 1\text{-butanol} \times \frac{0.8279 \text{ g } 1\text{-butanol}}{1 \text{ mL } 1\text{-butanol}} = 13.77 \text{ g } 1\text{-butanol}
\]

Mass of liquid from the density.

\[
13.77 \text{ g } 1\text{-butanol} \times \frac{1 \text{ mole } 1\text{-butanol}}{74.12 \text{ g } 1\text{-butanol}} = 0.1857 \text{ moles } 1\text{-butanol}
\]

Moles of starting material.

\[
0.1857 \text{ moles } 1\text{-butanol} \times \frac{1 \text{ mole } 1\text{-bromobutane}}{1 \text{ mole } 1\text{-butanol}} = 0.1857 \text{ moles } 1\text{-bromobutane}
\]

Moles of product.

\[
0.1857 \text{ moles } 1\text{-bromobutane} \times \frac{137.03 \text{ g } 1\text{-bromobutane}}{1 \text{ mole } 1\text{-bromobutane}} = 25.47 \text{ g } 1\text{-bromobutane}
\]

Grams of product (calculated). This is theoretical yield.

Apparatus: Reaction setup uses in Zuluich with gas (HBr) trap at top of reflux condenser:

- Air in
- Distilling adapter
- OPEN inlet adapter

Local modification duly noted.

To the precipitate with trap (slow flow) pulls vapors into water stream.

**Fig. 6** Notebook entry for a synthesis experiment (2).
Synthesis of 1-Bromobutane (cont'd)

Outline of Procedure

1. Place 24.0 g NaBr, 25 ml water, and 17 ml 1-butanol in a 250 ml R.B. flask and cool in ice-water bath to < 10°C.
2. SLOWLY add 20 ml conc. H$_2$SO$_4$ with swirling.
3. Reflux this mixture over a flame for 30 min.
4. Let mixture cool. Distill mixture - receiver cooled in ice-water bath. Distill until distillate is NOT cloudy.
5. Collect a few drops of clear distillate in a test tube. Add water and shake tube. If two layers form, continue distilling another 5-10 min. and repeat this test. If two layers do not form, distill for another 5-10 minute and quit.
6. Wash distillate with 25 ml H$_2$O.
7. Wash distillate with 15 ml cold conc. H$_2$SO$_4$.
8. Wash distillate with 15 ml 10% sodium hydroxide solution.
9. Dry with anhydrous MgSO$_4$ & filter (gravity) into small, dry R.B. flask.

Fig. 7 Notebook entry for a synthesis experiment (3).
Synthesis of 1-Bromotoluene (cont'd.)

Placed 240 g NaBr, 25 ml water and 17 ml 1-butanol in a 250 ml R.B. flask. Let cool in an ice-bath. When liquid reached 5°C, added H₂SO₄ with swirling. The mixture warmed up and turned a yellow color.

Let up for reflux with gas tap as on p. 27. Mixture darkened as reflux continued.

Placed 15 ml H₂SO₄ in eulynmeyer clamped to cool in ice-water bath for later.

After refluxing 30 min, let reaction mixture cool to room temp, then put in ice-water bath. There are two distinct layers on the flask, both an orange color. Color may be due to free bromine. One of the two layers is product.

Distilled the mixture, and collected everything that came over up to 100°C. Initially, cloudy, white liquid came over (water + organic product?) then clear liquid. Stopped heating, bubbly flask, removed receiver and replaced it with test-tube in beaker with 1°C water. Heated to distil over a few drops of liquid. Added a bit more than an equal amount of water shock tube. No layers formed! Replaced receiver, flask & distilled for 5 min, more.

Poured distillate into a 125 ml eulynmeyer, separating funnel; added 25 ml water. Water went into upper layer - upper layer as aqueous; lower as organic product.

Note the recorded observations.

Next step performed while reflux continues.

Only a phrase recalls the distillation.

Fig. 8 Notebook entry for a synthesis experiment (4).
Synthesis of 1 Bromotoluene (cont'd)

Washed product with the 15 ml cold conc. H2SO4. Solution warmed up as I shook flask!

Washed with the 15 ml 10% sodium hydroxide. Added 5 ml water and it stayed in upper layer. Treated aqueous layer with red litmus paper and it turned blue — so organic layer is not acidic.

Put product into 50 ml Erlenmeyer and added anhydrous magnesium sulfate in small amounts with swirling. Cloudy product turned clear and XS seemed drying agent was working on the flask. Cycled flask 4 times at away.

10/19/86 J.B.

10/14/86

Set up for distillation. Removed thermometer and thermometer adaptor and put long-stemmed funnel into the flask. Gravity filtered my product directly into distilling flask. Dropped in boiling stone, replaced thermometer of adaptor, and distilled legend. Collected all that came over from 100-103°C.

---

Fig. 9  Notebook entry for a synthesis experiment (5).
Fig. 10 Notebook entry for a synthesis experiment (6).

Synthesis of 1-Bromobutene (cont'd)

Weight of labelled vial 20.2 g

Weight

Weight of vial and product 36.6 g
Weight of labelled vial 20.2 g

wt. of product 16.2 g

Put product into clean, weighed, labelled vial.
Product yield 16.2 g

% yield = \( \frac{16.2 \text{ g}}{25.44 \text{ g}} \times 100 = 63.6 \% \)
present in the smallest molar ratio. Note the use of the density to get from volume to mass (ml to g), molecular weight to go from mass to number of moles (g to mol), the stoichiometric ratio (here 1:1) to get moles of product from moles of limiting reagent, and finally reapplication of molecular weight to get the mass (g) of the product. Note that this mass is calculated. It is NOT anything we've actually produced. In THEORY, we get this much. That is theoretical yield.

5. I'm a firm believer in the use of units, factor-label method, dimensional analysis, whatever you call it. I KNOW I've screwed up if my units are (g 1-butanol)²/mole 1-butanol.

6. Remember the huge writeup on the Separation of a Liquid Mixture by Distillation, drawings of apparatus and all? Well, the line “the mixture was purified by distillation,” (Fig. 9) is all you write for the distillation during this synthesis.

7. At the end of the synthesis, you calculate the percent yield. Just divide the amount you actually prepared by the amount you calculated you’d get, and multiply this fraction by 100. For this synthesis, I calculated a yield of 25.44 g of product. For this reaction on the bench, I actually obtained 16.2 g of product. So:

\[
(16.2 \text{ g} / 25.44 \text{ g})(100) = 63.6\% \text{ yield}
\]
You should look up information concerning any organic chemical you'll be working with so that you know what to expect in terms of molecular weight, density, solubility, crystalline form, melting or boiling point, color, and so on. This information is kept in handbooks that should be available in the lab, if not in the library. Reading some of these is not easy, but once someone tells you what some of the fancy symbols mean, there shouldn't be a problem. Many of the symbols are common to all handbooks, and are discussed only once, so read the entire section even if your handbook is different. There are at least four fairly popular handbooks and I've included sample entries of 1-bromobutane and benzoic acid, a liquid and a solid you might come across in lab, to help explain things.

**CRC HANDBOOK**

*(CRC Handbook of Chemistry and Physics, CRC Press, Inc., Boca Raton, Florida.)* Commonly called “the CRC” as in, “Look it up in the CRC.” A very popular book; a classic. Sometimes you can get the last year’s edition cheaply from the publisher, but it’s usually for an order of 10 or more.

**Entry: 1-Bromobutane (Fig. 11)**

1. **No. 3683.** An internal reference number. Other tables in the handbook will use this number, rather than the name.
2. **Name, . . . Butane, 1-bromo.** You get a systematic name and a formula.
3. **Mol. wt. 137.03.** The molecular weight of 1-bromobutane.
4. **Color, . . . . . . . Dots!** This implies 1-bromobutane is a colorless liquid; nothing special really.
5. **b. p. 101.6, 18.8°.** The normaling boiling point, at 760 torr, is 101.6°C. The 18.8 has a tiny superscript to tell you that 18.8°C is the boiling point at 30 torr.
6. **m.p. −112.4.** The melting point of solid 1-bromobutane. Handbooks report only the TOP of the melting point range. You, however, should report the entire range.
7. **Density. 1.2758\(^{20/4}\).** Actually, this particular number is a specific gravity. This is a mass of the density of the liquid taken at 20°C referred to (divided by) the density of the same mass of water at 4°C. That’s what the tiny 20/4 means. Notice the units will cancel. A number without the modifying fraction is a true density (in g/ml) at the temperature given.

8. **\(n_D 1.4401^{20}\).** This is the index of refraction (see Chapter 22, “Refractometry”) obtained using the yellow light from a sodium lamp (the D line). Yes, the tiny 20 means it was taken at 20°C.

9. **Solubility. al, eth, ace, chl.** This is what 1-bromobutane must be soluble in. There are a lot of solvents, and here are the abbreviations for some of them:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>al</td>
<td>alcohol</td>
</tr>
<tr>
<td>bz</td>
<td>benzene</td>
</tr>
<tr>
<td>peth</td>
<td>petroleum ether</td>
</tr>
<tr>
<td>aa</td>
<td>acetic acid</td>
</tr>
<tr>
<td>lig</td>
<td>ligroin</td>
</tr>
<tr>
<td>to</td>
<td>toluene</td>
</tr>
<tr>
<td>eth</td>
<td>ether</td>
</tr>
<tr>
<td>chl</td>
<td>chloroform</td>
</tr>
<tr>
<td>w</td>
<td>water</td>
</tr>
<tr>
<td>MeOH</td>
<td>methanol</td>
</tr>
<tr>
<td>CCl(_4)</td>
<td>carbon tetrachloride</td>
</tr>
</tbody>
</table>

Some solvents have such a long tradition of use, they are our old friends and we use very informal names for them:

- **alcohol.** Ethyl alcohol; ethanol.
- **ether.** Diethyl ether; ethoxyethane.
- **pet. ether.** Petroleum ether. Not a true ether, but a low boiling (30–60°C) hydrocarbon fraction like gasoline.
- **ligroin.** Another hydrocarbon mixture with a higher boiling range (60–90°C) than pet. ether.

10. **Ref. B1\(^4\) 258** Reference to listing in a set of German handbooks called “Beilstein.” Pronounce the German “ei” like the long i and stop yourself from saying “Beelsteen” or some such nonsense. 1-Bromobutane is in the fourth supplement (\(^4\)), Volume 1 (B1) on page 258.
<table>
<thead>
<tr>
<th>No.</th>
<th>Name, Synonyms, and Formula</th>
<th>Mol. wt.</th>
<th>Color, crystalline form, specific rotation and ( \lambda_{max} ) (log ( \epsilon ))</th>
<th>b.p. °C</th>
<th>m.p. °C</th>
<th>Density</th>
<th>nD</th>
<th>Solubility</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2531</td>
<td>Benzoic acid</td>
<td>122.13</td>
<td>mcl if or nd</td>
<td>249,133</td>
<td>122.13</td>
<td>1.0749(^{130}) 1.2659(^{13/4})</td>
<td>1.504(^{12})</td>
<td>al, eth, ace, bz, chl</td>
<td>B9(^{3}), 360</td>
</tr>
<tr>
<td>2532</td>
<td>Benzoic acid, 2-acetamido</td>
<td>179.18</td>
<td>nd (aa)</td>
<td></td>
<td>185</td>
<td></td>
<td></td>
<td>eth, ace</td>
<td>B14(^{3}), 922</td>
</tr>
<tr>
<td>3683</td>
<td>Butane, 1-bromo</td>
<td>137.03</td>
<td></td>
<td>101.6, 18.8(^{30})</td>
<td>−112.4</td>
<td>1.2758(^{20/4}) 1.440(^{12})</td>
<td>al, eth, ace, chl</td>
<td>B14(^{4}), 258</td>
<td></td>
</tr>
<tr>
<td>3684</td>
<td>Butane, 1-bromo-4-chloro</td>
<td>171.48</td>
<td></td>
<td>174−5(^{38}), 63−4(^{10})</td>
<td></td>
<td>1.488(^{20/4}) 1.4885(^{20})</td>
<td>al, eth, chl</td>
<td>B14(^{4}), 264</td>
<td></td>
</tr>
</tbody>
</table>

**Fig. 11** Sample CRC entries from the 61st edition.
Entry: Benzoic Acid (Fig. 11)

There are a few differences in the entries, what with benzoic acid being a solid, and I'll point these out. If I don't reexamine a heading, see the explanation back in the 1-Bromobutane entry for details.

1. **Color, . . . mcl lf or nd.** Monoclinic leaflets or needles. This is the shape of the crystals. There are many different crystalline shapes and colors and I can't list them all—but here's a few:

   pl plates  mcl monoclinic
   nd needles  rh rhombus
   lf leaves  ye yellow
   pr prisms  pa pale

   I've included #2532 (Benzoic acid, 2-acetamido) to show that you sometimes get a bonus. Here nd(aa) means you get needle-like crystals from acetic acid. Acetic acid (aa) is the recrystallization solvent (see Chapter 10, "Recrystallization"), and you don't have to find it on your own. Thus, pa ye nd (al) means that pale yellow needles are obtained when you recrystallize the compound from ethanol.

2. **Density. 1.0749**. This is an actual density of benzoic acid taken at 130°C. There is no temperature ratio as there is for the specific gravity (1.2659$^{15/4}$).

Nostalgia

I've included the entries from the 43rd and 49th editions of the CRC to show you that not all things improve with age.

1. **General Organization.** The 43rd and 49th editions make use of **bold-face type** to list parent compounds, and lighter type to list derivatives. Benzoic acid is a parent; there are many derivatives (Fig. 12). The 61st edition lists all compounds with the same weight (Fig. 11).

2. **Solubility tables.** Here the older editions really shine. The 43rd edition gives numerical solubility data for benzoic acid: 0.18$^4$, 0.27$^{18}$, 2.2$^{75}$. These
### PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS (Continued)

<table>
<thead>
<tr>
<th></th>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>b1239</td>
<td>Benzolic acid</td>
<td>Benzenecarboxylic acid.</td>
<td>122.12</td>
<td>mcl if or nd</td>
<td>122.4</td>
<td>249(^{780})</td>
<td>1.3211(^{233})</td>
<td>1.504(^{132})</td>
<td>δ</td>
<td>v</td>
</tr>
<tr>
<td>b1240</td>
<td>—, 4-acetamidophenyl ester</td>
<td>p-Benzoyloxyacetanilide. C(\text{H}_4)(\text{NO}_2), See b1239</td>
<td>255.28</td>
<td>nd (al)</td>
<td>171</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>b2500</td>
<td>Butane</td>
<td>—, 1-bromo-*</td>
<td>137.03</td>
<td>—</td>
<td>-112.3</td>
<td>101.3(^{780})</td>
<td>1.2764(^{20})</td>
<td>1.4398(^{20})</td>
<td>1</td>
<td>∞</td>
</tr>
</tbody>
</table>

**Fig. 12** Sample CRC entries from the 49th edition.
are the actual solubilities, in grams of benzoic acid per 100 g of water at 4, 18, and 75°C, respectively. The butyl bromide (1-bromobutane) entry has helpful solubility indicators: i, insoluble in water; ∞, miscible in alcohol; ∞, miscible in diethyl ether.

There are other abbreviations used for the solubility of a compound. Some of the more popular abbreviations are

<table>
<thead>
<tr>
<th>s</th>
<th>soluble</th>
<th>i</th>
<th>insoluble</th>
</tr>
</thead>
<tbody>
<tr>
<td>δ</td>
<td>slightly soluble</td>
<td>∞</td>
<td>miscible, mixes in all proportions.</td>
</tr>
<tr>
<td>h</td>
<td>solvent must be hot</td>
<td>v</td>
<td>very</td>
</tr>
</tbody>
</table>

What a big change from the 43rd to the 61st edition. Numerical solubility data missing, solubility indications gone, and even incomplete solubility reporting (Benzoic acid: chl, CCl₄, acet., me. al., bz, CS₂—43rd ed.; where are CCl₄, me. al., and CS₂ in the 61st?). The decrease in organizational structure, I can live with. But the new way of presenting the solubility data (what there is of it) is useless for many things you need to do in your lab. Reread the sample synthesis experiment (see Chapter 2, “Keeping a Notebook”). You need more useful solubility data for that experiment than you can extract from the most recent CRC Handbook. For my money, you want a fancy $17.50 doorstop, get a CRC 61st and up. You want useful information, get a CRC 60th and back. Or consult the handbook I want to talk about next.
### PHYSICAL CONSTANTS OF

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
<th>Synonyms</th>
<th>Formula</th>
<th>Mol. Wt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1449</td>
<td>Benzoic acid</td>
<td>benzene carboxylic acid*, phenylformic acid</td>
<td>C₆H₅COOH</td>
<td>122.12</td>
</tr>
<tr>
<td>1450</td>
<td>—, allyl ester</td>
<td>allyl benzoate</td>
<td>C₆H₅COOC₂H₅</td>
<td>162.18</td>
</tr>
<tr>
<td>1451</td>
<td>—, anhydride</td>
<td>See Benzoic anhydride.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1452</td>
<td>—, benzyl ester</td>
<td>benzyl benzoate; benzyl benzenecarboxylate</td>
<td>C₆H₅COOCH₂C₆H₅</td>
<td>212.24</td>
</tr>
</tbody>
</table>

*Name approved by the International Union of Chemistry.

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### PHYSICAL CONSTANTS OF

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
<th>Synonyms</th>
<th>Formula</th>
<th>Mol. Wt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2160</td>
<td>Butyl bromide (n)</td>
<td>1-bromobutane*</td>
<td>CH₃(CH₂)₂CH₂Br</td>
<td>137.03</td>
</tr>
<tr>
<td>2161</td>
<td>sec-Butyl bromide</td>
<td>2-bromobutane*, methyl-ethylbromomethane</td>
<td>C₃H₇CH(CH₃)Br</td>
<td>137.03</td>
</tr>
<tr>
<td>2162</td>
<td>tert-Butyl bromide</td>
<td>2-bromo-2-methylpropane*, trimethylbromomethane</td>
<td>(CH₃)₃CBr</td>
<td>137.03</td>
</tr>
</tbody>
</table>

*Name approved by the International Union of Chemistry.

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Fig. 13  Sample CRC entries from the 41st edition.

Entry: 1-Bromobutane (Fig. 14)

1. **Name. Butyl bromide(n).** Here, 1-bromobutane is listed as a substituted butyl group much like in the 43rd CRC. The systematic name is listed under synonyms.

2. **Beil. Ref. I-119.** The Beilstein reference; Volume 1, page 119, the original work (not a supplement).
### ORGANIC COMPOUNDS (Continued)

<table>
<thead>
<tr>
<th>No.</th>
<th>Crystalline form, color and index of refraction</th>
<th>Density g/ml</th>
<th>Melting point, °C</th>
<th>Boiling point, °C</th>
<th>Solubility in grams per 100 ml of Water</th>
<th>Alcohol</th>
<th>Ether, etc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1449</td>
<td>col. monoc. leaf. or need., 1.53974(^{15})</td>
<td>1.2659(^{1/2})</td>
<td>122</td>
<td>249</td>
<td>0.18(^{4})</td>
<td>0.27(^{18})</td>
<td>2.2(^{25})</td>
</tr>
<tr>
<td>1450</td>
<td>yel. liq.</td>
<td>1.058(^{1/2})</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1451</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1452</td>
<td>col. oily liq., or need. or leaf., 1.5681(^{21})</td>
<td>1.114(^{18})</td>
<td>21 (18.5)</td>
<td>323–4</td>
<td>00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For explanations and abbreviations see beginning of table.

### ORGANIC COMPOUNDS (Continued)

<table>
<thead>
<tr>
<th>No.</th>
<th>Crystalline form, color and index of refraction</th>
<th>Density g/ml</th>
<th>Melting point, °C</th>
<th>Boiling point, °C</th>
<th>Solubility in grams per 100 ml of Water</th>
<th>Alcohol</th>
<th>Ether, etc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2160</td>
<td>col. liq., 1.4398</td>
<td>1.299(^{20})</td>
<td>-112.4</td>
<td>101.6</td>
<td>i.</td>
<td>^(\infty)</td>
<td>^(\infty) eth.</td>
</tr>
<tr>
<td>2161</td>
<td>col. liq., 1.4344(^{25})</td>
<td>1.2580(^{20})</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2162</td>
<td>col, liq., 1.428</td>
<td>1.222(^{21})</td>
<td>-20</td>
<td>73.3</td>
<td>i.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For explanations and abbreviations see beginning of table.

---

3. **Crystalline form . . . lq.** It's a liquid.

4. **Specific gravity 1.275\(^{20}/4\).** The tiny temperature notation is presented a bit differently, but the meaning is the same.

5. **Solubility in 100 parts water. 0.06\(^{16}\)** 0.06g of 1-bromobutane will dissolve in 100 g of water at 16°C. After that, no more.
### Table 7-4 (Continued)

PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
<th>Synonym</th>
<th>Formula</th>
<th>Beil. Ref.</th>
<th>Formula Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>711</td>
<td>Benzoic acid</td>
<td></td>
<td>C₆H₅·CO₂H</td>
<td>IX-92</td>
<td>122.12</td>
</tr>
<tr>
<td>712</td>
<td>Na salt</td>
<td>sodium benzoate</td>
<td>C₆H₅·CO₂Na·H₂O</td>
<td>IX-107</td>
<td>162.12</td>
</tr>
</tbody>
</table>

Benznaphthalide 765  
Benzoic acid sulfamide 5671-3

### Table 7-4 (Continued)

PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
<th>Synonym</th>
<th>Formula</th>
<th>Beil. Ref.</th>
<th>Formula Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1040</td>
<td>Butyl amine (sec)</td>
<td></td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{NH})</td>
<td>IV-160</td>
<td>73.14</td>
</tr>
<tr>
<td>1041</td>
<td>amine (iso)</td>
<td></td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{NH}_2)</td>
<td>IV-163</td>
<td>73.14</td>
</tr>
<tr>
<td>1057</td>
<td>bromide (n)</td>
<td>1-bromo-butane</td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{Br})</td>
<td>I-119</td>
<td>137.03</td>
</tr>
<tr>
<td>1058</td>
<td>bromide (sec)</td>
<td>2-bromo-butane</td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{Br})</td>
<td>I-119</td>
<td>137.03</td>
</tr>
<tr>
<td>1059</td>
<td>bromide (iso)</td>
<td>1-Br-2-Me-propane</td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{Br})</td>
<td>I-126</td>
<td>137.03</td>
</tr>
<tr>
<td>1060</td>
<td>bromide ( tert)</td>
<td>2-Br-2-Me-propane</td>
<td>(\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{Br})</td>
<td>I-127</td>
<td>137.03</td>
</tr>
</tbody>
</table>

Butyl borate 6117  
Butyl carbamid 1138-9  
Butyl carbinol (iso) 406  
Butyl carbinol (sec) 411  
Butyl carbinol (n) 404  
Butyl carbinol ( tert) 410  
Butyl carbinol 2232

**Fig. 14** Sample entries from Lange's 11th edition.

**Entry: Benzoic Acid (Fig. 14)**

1. **Melting point. subl > 100.** Benzoic acid starts to sublime (go directly from a solid to a vapor) over 100°C, before any crystals left melt at 122.4°C.
### ORGANIC CHEMISTRY

**Table 7-4 (Continued)**

**PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS**

<table>
<thead>
<tr>
<th>No.</th>
<th>Crystalline Form and Color</th>
<th>Specific Gravity</th>
<th>Melting Point °C.</th>
<th>Boiling Point °C.</th>
<th>Solubility in 100 Parts</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Water</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Alcohol</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ether</td>
</tr>
<tr>
<td>711</td>
<td>mn. pr.</td>
<td>1.3163°</td>
<td>122.4; subl. &gt; 100</td>
<td>250.0</td>
<td>0.2117°; 61°; 2.2°</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.2176°; 2.2°; abs. al.</td>
</tr>
<tr>
<td>712</td>
<td>col. cr.</td>
<td>0.7242°</td>
<td>0.7323°</td>
<td>1.2752°</td>
<td>1.2612°</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.2642°</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.2203°</td>
</tr>
</tbody>
</table>

Benzophenone oxide 6451

### ORGANIC CHEMISTRY

**Table 7-4 (Continued)**

**PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS**

<table>
<thead>
<tr>
<th>No.</th>
<th>Crystalline Form and Color</th>
<th>Specific Gravity</th>
<th>Melting Point °C.</th>
<th>Boiling Point °C.</th>
<th>Solubility in 100 Parts</th>
</tr>
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<tr>
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<td></td>
<td></td>
<td>Water</td>
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<td></td>
<td></td>
<td>Alcohol</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ether</td>
</tr>
<tr>
<td>1040</td>
<td>col. iq.</td>
<td>0.7242°</td>
<td>-104</td>
<td>6677°mm</td>
<td>∞</td>
</tr>
<tr>
<td>1041</td>
<td>col. iq.</td>
<td>0.7323°</td>
<td>-85</td>
<td>68.9</td>
<td>∞</td>
</tr>
<tr>
<td>1057</td>
<td>liq.</td>
<td>1.2752°</td>
<td>-112.4</td>
<td>101.6</td>
<td>0.0618°</td>
</tr>
<tr>
<td>1058</td>
<td>liq.</td>
<td>1.2612°</td>
<td>-112.1</td>
<td>91.3</td>
<td>l.</td>
</tr>
<tr>
<td>1059</td>
<td>liq.</td>
<td>1.2642°</td>
<td>-117.4</td>
<td>91.4</td>
<td>0.0618°</td>
</tr>
<tr>
<td>1060</td>
<td>liq.</td>
<td>1.2203°</td>
<td>-16.2</td>
<td>73.3; d. 210</td>
<td>0.0618°</td>
</tr>
</tbody>
</table>

Butyl carbonate 1892-4
Butyl citrate 6119
Butyl cyanide (iso) 6413
Butyl cyanide (tort) 6246

2. **Crystalline form . . . , mn. pr.** monoclinic prisms. Here, *mn* is a variant of the *mcl* abbreviation used in the *CRC*. Don’t let these small differences throw you. A secret is that all handbooks have a listing of abbreviations at the front of the tables. Shhhh! Don’t tell anyone. It’s a secret.
I like the Lange’s format, redolent of the 43rd edition of the CRC. The one with the useful information. The organization based on common names, rather than systematic names, can make finding an entry a bit more difficult. There’s a miniature gloss at the bottom of each page to help you find related compounds.

Butyl carbinol(n), at the bottom of Fig. 14, has an index number of 404. If you’re familiar with the carbinol naming scheme for alcohols, it isn’t much to translate that to 1-pentanol. The entry still comes before the B’s because Amyl alcohol(n), is another common name for 1-pentanol. On the page where 1-pentanol would show up, there’s only a gloss entry: 1-Pentanol, 404. This brings you right back to Amyl alcohol(n). Since most textbooks and labbooks are making it a very big deal these days to list none but the purest of pristine systematic nomenclature, you’d likely never expect the compounds to be listed this way, and that is a bit annoying. Even though you are missing out on a bit of the history in the field.

MERCK INDEX

(The Merck Index, Merck & Co., Inc., Rahway, New Jersey.) This handbook is mostly concerned with drugs and their physiological effects. But useful information exists concerning many chemicals. Because of the nature of the listings, I’ve had to treat the explanations a bit differently than those for the other handbooks.

Entry: 1-Bromobutane (Fig. 15)

1. **Top of page. 1522 n-Butylbenzene.** Just like a dictionary, each page has headings directing you to the first entry on that page. So, 1522 is not the page number but the compound number for n-Butylbenzene, the first entry on page 216. The actual page number is at the bottom left of the page.

2. **n-Butyl Bromide.** Listed as a substituted butane with the systematic name given as a synonym.
3. \( C 35.06\% \). Elemental analysis data; the percent of each element in the compound.

4. *Prepd from*. A short note on how 1-bromobutane has been prepared, and references to the original literature (journals).

5. \( d_2^{25} 1.2686 \). The tiny 25 over 4 makes this a specific gravity. Note that the temperatures are given with the \( d \) and not with the numerical value as in Lange's and the CRC.

**Entry: Benzoic Acid (Fig. 15)**

1. *Line 2. dracylic acid*. What a synonym! Label your benzoic acid bottles this way and no one will ever "borrow" your benzoic acid again.


3. *Lines 7–9*. Industrial syntheses of benzoic acid. These are usually not appropriate for your lab bench preparations.

4. *Lines 9–20*. References to the preparation and characteristics of benzoic acid in the original literature (journals).


6. *Lines 21–40. Physical data*. The usual crystalline shape, density (note two values reported.), sublimation notation, boiling point data, and so on. \( K \) at 25° is the ionization constant of the acid; the pH of the saturated solution (2.8 at 25° C) is given. The solubility data (Soly) is very complete, including water solutions at various temperatures, a bit about the phase diagram of the compound, and solubility in other solvents. Note that numerical data is given where possible.


10. *Lines 73–75*. Therapeutic uses, both human and veterinary, for benzoic acid.

If all the chemical entries were as extensive as the one for benzoic acid, this would be the handbook of choice. Because benzoic acid has wide use in medi-
1522

\[ n \text{-Butylbenzene} \]

1526. \( n \)-Butyl Bromide. \( 1 \)-Bromobutane. \( C_4H_9Br \); mol wt 137.03. C 35.06%, H 6.62%, Br 58.32%. \( CH_2(CH_2)Br \). Prepd from \( n \)-butyl alc and a hydrobromic-sulfuric acid mixture: Kamm, Marvel, \textit{Org. Syn.} vol. 1, 5 (1921); Skau, McCullough, \textit{J. Am. Chem. Soc.} 57, 2440 (1935).

Colorless liquid. \( d_2^0=1.2686 \). \( bp_{760} \) 101.3° (mp -112°). \( n^0_2=1.4398 \). Insol in water; sol in alcohol, ether.

Page 216 Consult the cross index before using this section.


![COOH](image)

Monoclinic tablets, plates, leaflets. d 1.321 (also reported as 1.266). mp 122.4°. Begins to sublime at around 100°. \( bp_{760} \) 249.2°; \( bp_{500} \) 277°; \( bp_{200} \) 205.8°; \( bp_{100} \) 186.2°; \( bp_{50} \) 172.8°; \( bp_{40} \) 162.6°; \( bp_{20} \) 146.7°; \( bp_{10} \) 132.1°. Volatile with steam. Flash pt 121-131°. K at 25°: 6.40 \( \times \) 10^-5; pH of satd soln at 25°: 2.8. Soly in water (g/l) at 0° = 1.7; at 10° = 2.1;

\textit{Fig. 15} Sample entries from the Merck Index, 10th edition.

cine and food production, and it is very important to know the physical properties of drugs and food additives, a lot of information on benzoic acid winds up in the \textit{Index}. 1-Bromobutane has little such use, and the size of the entry reflects this. Unfortunately, many of the compounds you come in contact with in the organic laboratory are going to be listed with about the same amount of information you’d find for 1-bromobutane, and not with the large quantities of data you’d find with benzoic acid.
at 20° = 2.9; at 25° = 3.4; at 30° = 4.2; at 40° = 6.0; at 50° = 9.5; at 60° = 12.0; at 70° = 17.7; at 80° = 27.5; at 90° = 45.5; at 95° = 68.0.
M'ixtures of excess benzoic acid and water form two liquid phases beginning at 89.7°. The two liquid phases unite at the critical soln temp of 117.2°. Composition of critical mixture: 32.34% benzoic acid, 67.66% water: see Ward, Cooper, J. Phys. Chem. 34, 1484 (1930). One gram dissolves in 2.3 ml cold alc, 1.5 ml boiling alc, 4.5 ml chloroform, 3 ml ether, 3 ml acetone, 30 ml carbon tetrachloride, 10 ml benzene, 30 ml carbon disulfide, 23 ml oil of turpentine; also sol in volatile and fixed oils, slightly in petr ether. The soly in water is increased by alkaline substances, such as borax or trisodium phosphate, see also Sodium Benzoate.

Barium salt dihydrate, \( \text{C}_{12}\text{H}_{10}\text{BaO}_4\cdot 2\text{H}_2\text{O} \), barium benzoate. Nacreous leaflets. Poisonous! Soluble in about 20 parts water; slightly sol in alc.

very sol in boiling water.
Calcium salt trihydrate, \( \text{C}_{10}\text{H}_{10}\text{CaO}_4\cdot 3\text{H}_2\text{O} \), calcium benzoate. Orthorhombic crystals or powder. d 1.44. Soluble in 25 parts water;

Cerium salt trihydrate, \( \text{C}_{2}\text{H}_{14}\text{CeO}_4\cdot 3\text{H}_2\text{O} \), cerous benzoate. White to reddish-white powder. Sol in hot water or hot alc.

Copper salt dihydrate, \( \text{C}_{12}\text{H}_{10}\text{CuO}_4\cdot 2\text{H}_2\text{O} \), cupric benzoate. Light blue, crys powder. Slightly soluble in cold water, more in hot water; sol in alc or in dil acids with separation of benzoic acid.

Lead salt dihydrate, \( \text{C}_{12}\text{H}_{10}\text{O}_4\cdot 2\text{H}_2\text{O} \), lead benzoate. Cryst powder. Poisonous! Slightly sol in water.

Manganese salt tetrahydrate, \( \text{C}_{12}\text{H}_{10}\text{MnO}_4\cdot 4\text{H}_2\text{O} \), manganese benzoate. Pale-red powder. Sol in water, alc. Also occurs with 3H₂O.

Nickel salt trihydrate, \( \text{C}_{12}\text{H}_{10}\text{NiO}_4\cdot 3\text{H}_2\text{O} \), nickel benzoate. Light-green odorless powder. Slightly sol in water; sol in ammonia; dec by acids.

Potassium salt trihydrate, \( \text{C}_{12}\text{H}_{10}\text{K}2 \cdot 3\text{H}_2\text{O} \), potassium benzoate. Cryst powder. Sol in water, alc.

Silversalt, \( \text{C}_{12}\text{H}_{10}\text{AgO}_2 \), silver benzoate. Light-sensitive powder. Sol in 385 parts cold water, more sol in hot water; very slightly sol in alc.

Uranium salt, \( \text{C}_{12}\text{H}_{10}\text{O}_2\text{U} \), uranum benzoate, uranyl benzoate. Yellow powder. Slightly sol in water, alc.

Toxicity: Mild irritant to skin, eyes, mucous membranes.

Use: Preserving foods, fats, fruit juices, alkaloidal solns, etc: manuf benzoates and benzoic acid comds, dye; as a mordant in calico printing: for curing tobacco. As standard in volumetric and calorimetric analysis.

Therap Cat: Pharmacutie aid (antifungal agent).
Therap Cat (vet): Has been used with salicylic acid as a topical antifungal.

THE ALDRICH CATALOG

(The Aldrich Catalog. Aldrich Chemical Co., Inc., Milwaukee, Wisconsin.)

Not your traditional hard-bound reference handbook, but a handy book, nonetheless. The company makes many compounds, some not yet listed in the other handbooks, and often gives structures and physical constants for them. As Aldrich is in the business of selling chemicals to industry, many industrial references are given.
Entry: 1-Bromobutane (Fig. 16)

1. **1-Bromobutane.** Here it is listed strictly alphabetically as it is — with all the bromo-compounds — not as a butane, 1-bromo-, and only a cross reference as a butyl bromide.

2. **[109-65-9].** This is the Chemical Abstracts Service (CAS) Registry number. *Chemical Abstracts*, published by the American Chemical Society, is a listing of the abstract or summary written for any paper in the chemical literature. Every compound made gets a number. This makes for easy searching by computer, as well as by hand.

3. **bp 100-104°.** Without a tiny superscript this is the boiling point at 760 torr.

4. **$n_D^{20} 1.4390.** Index of refraction. The temperature (20°) modifies the $n$, rather than the number as in the CRC.

5. **d 1.276.** The density in g/cc.

6. **Fp 75°F(23°C) Flash point.** Above 75°F, a mixture of 1-bromobutane and air and a spark will go up like gangbusters. Watch out!

7. **Beil. 1,119.** The Beilstein reference; Volume 1, page 119.

8. **Merck Index 10, 1526.** The Merck Index 10th ed. reference; compound #1526 (Fig. 15).


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**Sample entries from the Aldrich catalog, 1986–87.**
10. **RTECS # EJ6225000.** The reference number in the Registry of Toxic Effects of Chemical Substances (RTECS). 1-Bromobutane is on the inventory of the EPA according to the Toxic Substances Control Act, PL94-69, October 11, 1976 (TSCA).

11. **Disp D.** There are methods of disposal given in the Aldrich Catalog. Go to method D and throw 1-bromobutane out according to the rules. Remember, the methods given are for the disposal of large amounts of a single substance, as might be found in an industrial application. The rules for the disposal of the waste generated in your undergraduate laboratory may differ considerably.

12. **FLAMMABLE LIQUID IRRITANT.** Yep, it sure is.

Note the differences in prices for the 99+% GOLD LABEL and the merely 99% 1-bromobutane. Before you buy, check on the use of the chemical. Normally, you can buy the least expensive grade of the chemical, and distill or recrystallize it yourself before you use it, if necessary.

**Entry: Benzoic Acid (Fig. 16)**

1. **Fieser 1, 49.** A reference to Fieser & Fieser's *Reagents for Organic Synthesis*, Volume 1, page 49. This multivolume series gives syntheses and reactions of many organic compounds, along with references to the original literature.

2. †. Benzoic acid cannot be shipped by parcel post.

3. **Beil. 9, 92.** A reference to Beilstein, Volume 9, page 92.


**NOT CLEAR—CLEAR?**

One antonym for **clear** is **cloudy**. Another antonym for **clear** is **colored**. When you say you “obtained a clear liquid,” do you mean that it is not cloudy, or that it is colorless?
Cloudiness usually means you’ve gotten water in your organic liquid. Colorless should be self-explanatory. You should always pair the turbidity and color designations:

"a clear, colorless liquid."
"a clear, blue liquid."
"a cloudy, colorless liquid."
"a cloudy, blue liquid."

I use clear to mean not cloudy, and *water-white* to mean not colored. Water-white is a designation found in the older chemical literature; *colorless* is more modern.

Is that clear?
Using **standard taper jointware** you can connect glassware without rubber stoppers, corks, or tubing. Pieces are joined by glass connections built into the apparatus (Fig. 17). They are manufactured in standard sizes, and you’ll probably use Т19/22.

The symbol Т means **standard taper**. The first number is the size of the joint at the widest point, in millimeters. The second number is the length of the joint, in millimeters. This is simple enough. Unfortunately, life is not all that simple, except for the mind that thought up this next devious little trick.

**STOPPERS WITH ONLY ONE NUMBER**

Sounds crazy, no? But with a very little imagination, and even less thought, grave problems can arise from confusing the two. Look at Fig. 18, which shows all glass stoppers are not alike. Interchanging these two leads to **leaking joints** through which your **graded** product can escape. Also, the Т19/22 stopper is much more expensive than the Т19 stopper, and you may have to

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**Fig. 17** Standard taper joints (T 19/22).
pay money to get the correct one when you check out at the end of the course. Please note the emphasis in those last two sentences. I appeal to your better nature and common sense. So, take some time to check these things out.

As you can see from Fig. 18, that single number is the width of the stopper at its top. There is no mention of the length, and you can see that it is too short. The T19 stopper does not fit the T19/22 joint. Only the T19/22 stopper can fit the T19/22 joint. Single-number stoppers are commonly used with volumetric flasks. Again, they will leak or stick if you put them in a double-number joint.

With these delightful words of warning, we continue the saga of coping with ground-glass jointware. Fig. 19 shows some of the more familiar pieces of jointware you may encounter in your travels. They may not be so familiar to you now, but give it time. After a semester or so, you’ll be good friends, go to reactions together, maybe take in a good synthesis. Real fun stuff!

These pieces of jointware are the more common pieces that I’ve seen used in the laboratory. You may or may not have all the pieces shown in Fig. 19. Nor will they necessarily be called EXACTLY by the names given here. The point is find out what each piece is, and make sure that it is in good condition before you sign your life away for it.
JOINTWARE

Vacuum adapter

Separator funnel

$\equiv$ Stopper

Thermometer (inlet) adapter

Round-bottom flasks

(Three-neck flask)

**Fig. 19** Some jointware.

ANOTHER EPISODE OF LOVE OF LABORATORY

"And that's $28.46 you owe us for the separatory funnel."

"But it was broken when I got it!"

"Should've reported it then."
"The guy at the next bench said it was only a two-dollar powder funnel and not to worry and the line at the stockroom was long anyway, and ... and ... anyway the stem was only cracked a little ... and it worked O.K. all year long ... Nobody said anything. ..."

"Sorry."

Tales like these are commonplace, and ignorance is no excuse. Don't rely on expert testimony from the person at the next bench. He may be more confused than you are. And equipment that is "slightly cracked" is much like a person who is "slightly dead." There is no in-between. If you are told that you must work with damaged equipment because there is no replacement available, you would do well to get it in writing.
HALL OF BLUNDERS AND THINGS NOT QUITE RIGHT

Round-Bottom Flasks

Round-bottom (R.B.) jointware flasks are so round and innocent looking, that you would never suspect they can turn on you in an instant.

1. **Star cracks.** A little talked about phenomenon that turns an ordinary R.B. flask into a potentially explosive monster. Stress, whether prolonged heating in one spot, or indiscriminate trouncing upon hard surfaces, can cause a flask to develop a **star crack** (Fig. 20) on its backside. Sometimes they are hard to see, but if overlooked, the flask can split asunder at the next lab.

2. **Heating a flask.** Since they are cold-blooded creatures, flasks show more of their unusual behavior while being heated. The behavior is usually unpleasant if certain precautions are not taken. In addition to star cracks, various states of disrepair can occur, leaving you with a benchtop to clean. Both humane and cruel heat treatment of flasks will be covered in (see Chapter 13, “Sources of Heat”), which is on the SPCG (Society for the Prevention of Cruelty to Glassware) recommended readings list.

![Fig. 20](image) R. B. flask with star crack.
Columns and Condensers

A word about **distilling columns** and **condensers**:

**Different!**

Use the **condenser** as is for **distillation** and **reflux** (see Chapter 15, "Distillation," and Chapter 16, "Reflux"). You can use the **column with or without column packing** (bits of metal or glass or ceramic or stainless-steel sponge — whatever)! That’s why the column is wider and it has **projections** at the end (Fig. 21). These projections help hold up the column packing if you use any packing at all (see Fig. 80).

If you jam column packing into the skinny condenser, the packing may never come out again! Using a condenser for a packed column is bad form and can lower your esteem or grade, whichever comes first.

**You might use the column as a condenser.**

**Never use the condenser as a packed column!**

The Adapter with Lots of Names

Fig. 22 shows the one place where joint and nonjoint apparatus meet. There are two parts: a rubber cap with a hole in it and a glass body. Think of the

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**Fig. 21** Distilling column versus condenser.
Rubber cap as a rubber stopper through which you can insert thermometers, inlet adapters, drying tubes, and so on.

**CAUTION!** Do not force. You might snap the part you’re trying to insert. Handle both pieces through a cloth; lubricate (water) and then insert carefully.

The rubber cap fits over the nonjoint end of the glass body. The other end is a *ground glass joint* and *fits only other glass joints*. The rubber cap should
neither crumble in your hands nor need a 10-ton press to bend it. If the cap is shot, get a new one. Let's have none of these corks, rubber stoppers, chewing gum, or any other type of plain vanilla adapter you may have hiding in the drawer.

*And remember:* Not only thermometers, but anything that resembles a glass tube can fit in here! This includes unlikely items such as drying tubes (they have an outlet tube) and even a funnel stem (you may have to couple the stem to a smaller glass tube if the stem is too fat).

The imaginative arrangements shown in Fig. 23 are acceptable.

**Forgetting the Glass**

Look, the Corning people went to a lot of trouble to turn out a piece of glass (Fig. 24) that fits perfectly in both a glass joint and a rubber adapter, so use it!

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**Fig. 23** Unusual, yet proper uses of the adapter with lots of names.
THINGS NOT TO DO WITH
THE ADAPTER WITH LOTS OF NAMES

**Fig. 24** The glassless glass adapter.

**Inserting Adapter Upside Down**

This one (Fig. 25) is really ingenious. If you're tempted in this direction, go sit in the corner and repeat over and over,

"Only glass joints fit glass joints"

**Fig. 25** The adapter stands on its head.
Inserting Adapter Upside Down *Sans* Glass
I don’t know whether to relate this problem (Fig. 26) to glass forgetting, or upside-downness, since it is both. Help me out. If I don’t see you trying to use an adapter upside down without the glass, I won’t have to make such a decision. So, don’t do it.

**GREASING THE JOINTS**

In all my time as an instructor, I’ve never had my students go overboard on greasing the joints, and they never got them stuck. Just lucky, I guess. Some instructors, however, use grease with a passion, and raise the roof over it. The entire concept of greasing joints is not as slippery as it may seem.

**To Grease or Not To Grease**

Generally you’ll grease joints on two occasions. One, when doing vacuum work to make a tight seal that can be undone; the other, doing reactions with
strong base that can etch the joints. Normally you don’t have to protect the joints during acid or neutral reactions.

**Preparation of the Joints**

Chances are you’ve inherited a set of jointware coated with 47 semesters of grease. First wipe off any grease with a towel. Then soak a rag in any hydrocarbon solvent (hexane, ligroin, petroleum ether — and no flames, these burn like gasoline) and wipe the joint again. Wash off any remaining grease with a strong soap solution. You may have to repeat the hydrocarbon–soap treatments to get a clean, grease-free joint.

**Into the Grease Pit (Fig. 27)**

First, use only enough to do the job! Spread it thinly along the upper part of the joints, only. Push the joints together with a twisting motion. The joint should turn clear from one third to one half of the way down the joint. *At no time should the entire joint clear!* This means you have *too much grease* and must start back at *Preparation of the Joints.*

Don’t interrupt the clear band around the joint. This is called *uneven greasing* and will cause you headaches later on.

**STORING STUFF AND STICKING STOPPERS**

At the end of a grueling lab session, you’re naturally anxious to leave. The reaction mixture is sitting in the joint flask, all through reacting for the day, waiting in anticipation for the next lab. You put the correct glass stopper in the flask, clean up, and leave.

*The next time, the stopper is stuck!*

Stuck but good! And you can probably kiss your flask, stopper, product and grade goodbye!
Frozen!

Some material has gotten into the glass joint seal, dried out, and cemented the flask shut. There are a few good cures, but several excellent preventive medicines.

Corks!

Yes, corks. Old-fashioned, non-stick-in-the-joint corks.
If the material you have to store does not attack cork, this is the cheapest, cleanest method of closing off a flask.
A well-greased glass stopper can be used for materials that attack cork, but
only if the stopper has a good coating of stopcock grease. Unfortunately, this grease can get into your product.

**Do not use rubber stoppers!**

Organic liquids can make rubber stoppers swell up like beach balls. The rubber dissolves and ruins your product, and the stopper won’t come out either. Ever.

The point is

**Dismantle all ground glass joints before you leave!**

**Corking a Vessel**

If winemakers corked their bottles like some people cork their flasks, there’d be few oneophiles and we’d probably judge good years for salad dressings rather than wines. You don’t just take a new cork and stick it down into the neck of the flask, vial, or what have you. You must press the cork first. Then as it expands, it makes a very good seal and doesn’t pop off.

A brand new cork, **before pressing** or **rolling**, should fit only about one-quarter of the way into the neck of the flask or vial. Then you roll the lower half of the cork on your clean benchtop to soften and press the small end. **Now** stopper your container. The cork will slowly expand a bit and make a very tight seal (Fig. 28).

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**Fig. 28** Corking a vessel.
Rather than rolling the cork on the benchtop, you might have the use of a **cork press**. You put the small end of the cork into the curved jaws of the press, and when you push the lever up and down, the grooved wheel rolls and mashes the cork at the same time (Fig. 29). Mind your fingers!
Other
Interesting
Equipment
An early edition of this book illustrated some equipment specific to the State University of New York at Buffalo, since that's where I was when I wrote it. It's now a few years later, and I realize that you can't make a comprehensive list.

Buffalo has an unusual "pear-shaped distilling flask" that I've not seen elsewhere. The University of Connecticut equipment list contains a "Bobbitt Filter Clip" that few other schools have picked up.

So if you are disappointed that I don't have a list and drawing of every single piece of equipment in your drawer, I apologize. Only the most common organic lab equipment is covered here. Ask your instructor "Whattizzit?" if you do not know.

I assume that you remember Erlenmeyer flasks, and beakers and such from the freshman lab. I'll discuss the other apparatus as it comes up in the various techniques. This might force you to read this book before you start lab.

Check out Fig. 30. Not all the mysterious doodads in your laboratory drawer are shown, but the more important are.
Fig. 30 Some stuff from your lab drawer.
Clean and Dry
Once you’ve identified your apparatus, you may find you have to clean it.

1. Wash your glassware at the end of the lab day. That way you’ll have clean and dry glassware, ready to go for the next lab. This may be difficult to do if you perform an experiment on the day you check in.

2. A little solvent, a little detergent, and a lot of elbow grease. These are the correct proportions for a cleaning solution. You do not need all the soap on the planet, nor do you have to fill the glassware to overflowing with soap solution. Agitation is the key here. The more you agitate a small amount of soap solution, the less you agitate your instructor by wasting your time and supplies, and the more effective your cleaning will be.

3. Special Buchner funnel cleaning alert. The standard ceramic Buchner funnel is not transparent, and you can’t see whether or not the bums who used the funnel the last time to collect a highly colored product, didn’t clean the funnel properly. The first time you Buchner filter crystals from an alcohol solution, the colored impurity dissolves, bleeds up into your previously clean crystals, and you may have to redo your entire experiment. I’d rinse the Buchner funnel with a bit of hot ethanol before I used it, just for insurance.

**Drying your glassware when you don’t need to**

“**It’s late. Why haven’t you started the experiment yet?”**

“I washed all my glassware and spent half an hour drying it.”

“What technique are we doing?”

“Steam distillation.”

“Steam goes through the entire setup, does it?”

A nodding head responds.

“What’s condensed steam?”

“Water. . . .”

There are all sorts of variations, but they boil down to this: You’ve taken all this time to dry your glassware only to put water in it. Writers of lab manuals are very tricky about this. Perhaps they say you’ll be using steam. Or maybe 5% aqueous sodium bicarbonate solution. Or even that a byproduct of your reaction is H₂O. Condense steam and you get what? An aqueous solution has what for a solvent? H₂O is what?
Look for sources of water other than plain water. If a "water-and" mixture is going to be in the equipment anyway, drying to perfection is silly.

If you wash your glassware before you quit for the day, the next time you need it, it'll be clean and dry. There are only a few reactions you might do that need superclean, superdry apparatus, and you should be given special instructions when that's necessary. (In their new book, Experimental Organic Chemistry, 2nd. Edition, McGraw-Hill, 1986, authors H. D. Durst and G. W. Gokel make the claim that glassware dried overnight is dry enough for the Grignard reaction, an extremely moisture-sensitive reaction, and flame drying can be avoided unless the laboratory atmosphere is extremely humid.)

Don't use the compressed air from the compressed air lines in the lab for drying anything. These systems are full of dirt, oil, and moisture from the pumps, and will get your equipment dirtier than before you washed it.

Yes, there are a few quick ways of drying glassware in case of emergency. You can rinse very wet glassware with a small amount of acetone, drain the glassware very well, and put the glassware in a drying oven (about 100°C) for a short spell. The acetone not only washes the water off the glassware very well (the two liquids are miscible, that is, they mix in all proportions.), the liquid left behind is acetone-rich, and evaporates faster than water. Don't use this technique unless absolutely necessary.
When you've prepared a liquid product, you must dry the liquid before you finally distill and package it, by treating the liquid with a **drying agent**. Drying agents are usually certain anhydrous salts that combine with the water in your product and hold it as a **water of crystallization**. When all of the water in your sample is tied up with the salt, you gravity filter the mixture. The dried liquid passes through the filter paper and the **hydrated salt** stays behind.

**TYPICAL DRYING AGENTS**

1. **Anhydrous calcium chloride.** This is a very popular drying agent, inexpensive and rapid, but of late I've become disappointed in its performance. It seems that the calcium chloride powders a bit upon storage and abuse, and this *calcium chloride dust* can go right through the filter paper with the liquid. So a caution: If you must use anhydrous calcium chloride, be sure it is granular. Avoid powdered calcium chloride, or granular anhydrous calcium chloride that's been around long enough to become pulverized. And don't add to the problem by leaving the lid off the jar of drying agent; that's the abuse I was talking about.

   Anhydrous calcium chloride tends to form *alcohols of crystallization*, so you really can't use it to dry alcohols.

2. **Anhydrous sodium carbonate and anhydrous potassium carbonate.** These are useful drying agents that are basically *basic*. As they dry your organic compound, any carbonate that gets dissolved in the tiny amounts of water in your sample can neutralize any tiny amounts of acid that may be left in the liquid. If your product is *supposed* to be acidic (in contrast to being *contaminated* with acid), you should avoid these drying agents.

3. **Anhydrous magnesium sulfate.** In my opinion, anh. MgSO₄ is about the best all-around drying agent. It has a drawback, though. Since it is a fine powder, lots of your product can become trapped on the surface. *This is not the same as water of crystallization.* The product is **only on the surface**, **not inside the crystal structure**, and you may wash your product off.
4. **Drierite.** Drierite, one commercially available brand of anhydrous calcium sulfate, has been around a long time and is a popular drying agent. You can put it in liquids and dry them or pack a drying tube with it to keep the moisture in the air from getting into the reaction setup. But be warned. There is also *Blue Drierite*. This has an indicator, a cobalt salt, that is *blue when dry, pink when wet*. Now you can easily tell when the drying agent is no good. Just look at it. Unfortunately, this stuff is not cheap, so don’t fill your entire drying tube with it just because it’ll look pretty. Use a small amount mixed with white Drierite, and when the blue pieces turn pink, change the entire charge in your drying tube. You can take a chance using Blue Drierite to dry a liquid directly. Sometimes the cobalt compound dissolves in your product. Then you have to clean and dry your product all over again.

**USING A DRYING AGENT**

1. Put the liquid or solution to be dried into an Erlenmeyer flask.
2. Add *small* amounts of drying agent and swirl the liquid. When the liquid is no longer cloudy, the water is gone, and the liquid is dry.
3. Add just a bit more drying agent and swirl one final time.
4. Gravity filter through filter paper (see Fig. 44).
5. If you’ve used a carrier solvent, then evaporate or distill it off, whichever is appropriate. Then you’ll have your clean, dry product.

**FOLLOWING DIRECTIONS AND LOSING PRODUCT ANYWAY**

“Add 5 g of anhydrous magnesium sulfate to dry the product.” Suppose your yield of product is lower than that in the book. Too much drying agent — not enough product — Zap! It’s all sucked onto the surface of the drying agent. Bye bye product. Bye bye grade.
Add the drying agent slowly to the product in small amounts

Now about those small amounts of product (usually liquids).

1. Dissolve your product in a low boiling point solvent. Maybe ether or hexane or the like. Now dry this whole solution, and gravity filter. Remove the solvent carefully. Hoo-ha! Dried product.

2. Use chunky dehydrating agents like anhydrous calcium sulfate (Drierite). Chunky drying agents have a much smaller surface area, so not much of the product gets adsorbed.
The fastest way to lose points is to hand in messy samples. Lots of things can happen to foul up your product. The following are unforgiveable sins!

**SOLID PRODUCTS**

1. **Trash in the sample.** Redissolve the sample, gravity filter, then evaporate the solvent.
2. **Wet solids.** Press out on filter paper, break up, let dry. The solid shouldn’t stick to the sides of the sample vial. Tacky!
3. **Extremely wet solids (solid floating in water).** Set up a gravity filtration (see “Gravity Filtration”) and filter the liquid off of the solid. Remove the filter paper cone with your solid product, open it up, and leave it to dry. Or remove the solid and dry it on fresh filter paper as above. Use lots of care though. You don’t want filter paper fibers trapped in your solid.

**LIQUID PRODUCTS**

1. **Water in the sample.** This shows up as droplets or as a layer of water on the top or the bottom of the vial, or *the sample is cloudy*. Dry the sample with a drying agent (see Chapter 7, “Drying Agents”) and gravity filter into a clean dry vial.
2. **Trash floating in the sample.** For that matter, it could be on the bottom, lying there. Gravity filter into a clean, dry vial.
3. **Water in the sample when you don’t have a lot of sample.** Since solid drying agents can absorb lots of liquid, what can you do if you have a tiny amount of product to be dried? Add some solvent that has a low boiling point. It must dissolve your product. Now you have a lot of liquid to dry, and *if a little gets lost, it is not all product*. Remove this solvent after you’ve dried the solution. Be careful if the solvent is flammable. *No flames!*
THE SAMPLE VIAL

Sad to say, but an attractive package can sell an inferior product. So why not sell yours. Dress it up in a neat new label. Put on

1. **Your name.** Just in case the sample gets lost on the way to camp.
2. **Product name.** So everyone will know what is in the vial. What does “Product from part C” mean to you? Nothing? Funny, it doesn’t mean anything to instructors either.
3. **Melting point (solids only).** This is a range, like “M.P. 96–98°C” (see Chapter 9, “The Melting Point Experiment”).
4. **Boiling point (liquids only).** This is a range “B.P. 96–98°C” (see Chapter 15, “Distillation”).
5. **Yield.** If you weigh the empty vial and cap, you have the tare weight. Now add your product and weigh the full vial. Subtract the tare weight from this gross weight to get the net weight (yield, in grams) of your product.
6. **Percent yield.** Calculate the percent yield (see Chapter 2, “Keeping a Notebook”) and put it on the label.

You may be asked for more data, but the things listed above are a good start down the road to good technique.

P.S. Gummed labels can fall off vials, and pencil will smear. *Always use waterproof ink!* And a piece of transparent tape over the label will keep it on.

HOLD IT! DON’T TOUCH THAT VIAL

Welcome to “You Bet Your Grade.” The secret word is **dissolve.** Say it slowly as you watch the cap liner in some vials dissolve into your nice, clean product and turn it all goopy. This can happen. A good way to prevent this is to cover the vial with aluminum foil before you put the cap on. Just make sure the product does not react with aluminum. Discuss this at length with your instructor.
The Melting Point Experiment
A **melting point** is the temperature at which the first crystal just starts to melt until the temperature at which the last crystal just disappears. Thus the melting point (abbreviated M.P.) is actually a **melting range**. You should report it as such, even though it is *called* a melting point, for example, M.P. 147–149°C.

People always read the phrase as melting *point* and never as melting *point*. There is this uncontrollable, driving urge to report one number. No matter how much I’ve screamed and shouted at people not to report one number, they almost always do. It’s probably because handbooks list only one number, the upper limit.

Generally, melting points are taken for two reasons.

1. **Determination of purity.** If you take a melting point of your compound and it starts melting at 60°C and doesn’t finish until 180°C you might suspect something is wrong. A melting range greater than 2°C usually indicates an impure compound (As with all rules, there are exceptions. There aren’t many to this one, though.).

2. **Identification of unknowns.**

   a. If you have an unknown solid, take a melting point. Many books (ask your instructor) contain tables of melting points and lists of compounds that may have a particular melting point. One of them may be your unknown. You may have 123 compounds to choose from. A little difficult, but that’s not all the compounds in the world. Who knows?? Give it a try. If nothing else, you know the melting point.

   b. Take your unknown and mix it *thoroughly* with some chemical you think might be your unknown. You might not get a sample of it, but you can ask. Shows you know something. Then:

   1) If the mixture melts at a *lower* temperature, over a *broad range*, your unknown is NOT the same compound.

   2) If the mixture melts at the *same temperature, same range*, it’s a good bet it’s the *same compound*. Try another one, though, with a different ratio of your unknown and this compound just to be sure. A *lower* melting point with a *sharp range* would be a special point called a **eutectic mixture**, and you, with all the other
troubles in lab, just might accidentally hit it. On lab quizzes, this is called

"Taking a mixed melting point."

Actually, "taking a mixture melting point," the melting point of a mixture, is more correct. But I have seen this expressed both ways.

**SAMPLE PREPARATION**

You usually take melting points in thin, closed end tubes called *capillary tubes*. They are also called *melting point tubes* or even *melting point capillaries*. The terms are interchangeable, and I'll use all three.

Sometimes you may get a supply of tubes that are open on both ends! You don't just use these as is. Light a burner, and close off one end, before you start. Otherwise your sample will fall out of the tube (see "Closing Off Melting Point Tubes," following).

Take melting points on *dry, solid* substances ONLY, *never* on liquids or solutions of solids in liquids or on wet or even damp solids.

*Only on dry solids!*

To help dry damp solids, place the damp solid on a piece of filter paper and fold the paper around the solid. Press. Repeat until the paper doesn't get wet. Yes, you may have to use fresh pieces of paper. Try not to get filter paper fibers in the sample, OK?

Occasionally, you may be tempted to dry solid samples in an oven. *Don't* — unless you are specifically instructed to. I know some students who have decomposed their products in ovens and under heat lamps. With the time they save quickly decomposing their product, they can repeat the entire experiment.

**Loading the Melting Point Tube**

Place a small amount of *dry solid* on a new filter paper (Fig. 31). Thrust the open end of the capillary tube into the middle of the pile of material. Some
THE MELTING POINT EXPERIMENT

solid should be trapped in the tube. Turn the tube over, closed end down. Remove any solid sticking to the outside. The solid must now be packed down.

Traditionally, the capillary tube, turned upright with the open end up, is stroked with a file, or tapped on the benchtop. Unless done carefully, these operations may break the tube. A safer method is to drop the tube closed end down, through a length of glass tubing. You can even use your condenser or distilling column for this purpose. When the capillary strikes the benchtop, the compound will be forced into the closed end. You may have to do this several times. If there is not enough material in the M.P. tube, thrust the open end of the tube into the mound of material and pack it down again. Use your own judgment; consult your instructor.

Use the smallest amount of material that can be seen to melt
Closing Off Melting Point Tubes

If you have melting point tubes that are *open at both ends* and you try to take a melting point with one, it should come as no surprise when your compound falls out of the tube. You’ll have to *close off one end*, to keep your sample from falling out (Fig. 32). So light a burner and get a “stiff” small blue flame. SLOWLY touch the end of the tube to the side of the flame, and hold it there. You should get a yellow sodium flame, and the tube will close up. There is no need to rotate the tube. And remember, *touch — just touch* — the edge of the flame, and hold the tube there. Don’t feel you have to push the tube way into the flame.

**MELTING POINT HINTS**

1. Use only the smallest amount that you can see melt. Larger samples will heat unevenly.
2. Pack down the material as much as you can. Left loose, the stuff will heat unevenly.
3. Never remelt any sample. They may undergo nasty chemical changes such as oxidation, rearrangement and decomposition.
4. Make up more than one sample. One is easy, two is easier. If something goes wrong with one, you have another. Duplicate, even triplicate runs are common.

Fig. 32  Closing off a M. P. tube with a flame.
THE MEL-TEMP APPARATUS

The Mel-Temp apparatus (Fig. 33) substitutes for the Thiele tube or open beaker and hot oil methods (see “Using the Thiele Tube”). Before you use the apparatus, there are a few things you should look for.

**Fig. 33** The Mel-Temp apparatus.
1. **Line cord.** Brings a.c. power to unit. Should be plugged into a live wall socket [See J. E. Leonard and L. E. Mohrmann, *J. Chem. Educ.*, 57, 119 (1980), for a modification in the wiring of older units, to make them less lethal. It seems that even with the three-prong plug, there can still be a shock hazard. *Make sure your instructor knows about this!*]

2. **On–off switch.** Turns the unit on or off.

3. **Fuse.** Provides electrical protection for the unit.

4. **Voltage control.** Controls the *rate of heating, not the temperature!* The higher the setting, the faster the temperature rise.

5. **Light source.** Provides illumination for samples.

6. **Eyepiece.** Magnifies the sample (Fig. 34).

7. **Thermometer.** Gives temperature of sample, and upsets the digestion when you’re not careful and you snap it off in the holder.

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**OPERATION OF THE MEL-TEMP APPARATUS**

1. *Imagine yourself getting burned if you’re not careful.* Never assume the unit is cold.

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*Fig. 34* Closeup of the viewing system.
2. Place loaded M.P. tube in one of the three channels in the opening at the top of the unit (Fig. 34).

3. Set the voltage control to zero if necessary. There are discourteous folk who do not reset the control when they finish using the equipment.

4. Turn the on-off switch to ON. The light source should illuminate the sample. If not, call for help.

5. Now science turns into art. Set the voltage control to any convenient setting. The point is to get up to within 20°C of the supposed melting point. Yep, that's right. If you have no idea what the melting point is, it may require several runs as you keep skipping past the point with a temperature rise of 5–10°C per minute. A convenient setting is 40. This is just a suggestion, not an article of faith.

6. After you've melted a sample, throw it away!

7. Once you have an idea of the melting point (or looked it up in a handbook, or were told), get a fresh sample, and bring the temperature up quickly at about 5–10°C per minute to within 20°C of this approximate melting point. Then turn down the voltage control to get a 2°C per minute rise. Patience!

8. When the first crystals just start to melt, record the temperature. When the last crystal just disappears, record the temperature. If both points appear to be the same, either the sample is extremely pure, or the temperature rise was too fast.

9. Turn the on-off switch to OFF. You can set the voltage control to zero for the next person.

10. Remove all capillary tubes.

Never use a wet rag or sponge to quickly cool off the heating block. This might permanently warp the block. You can use a cold metal block to cool it if you're in a hurry. Careful. If you slip, you may burn yourself.

THE FISHER–JOHNS APPARATUS

The Fisher–Johns apparatus (Fig. 35) is different in that you don’t use capillary tubes to hold the sample. Instead, you sandwich your sample be-
tween two round microscope cover slides (thin windows of glass) on a heating block. This type of melting point apparatus is called a **hot stage**. It comes complete with spotlight Look for the following.

1. **Line Cord** (at the back). Brings a.c. power to unit. Should be plugged into a live wall socket.
2. **On-off switch.** Turns the unit on or off.
3. **Fuse** (also at the back). Provides electrical protection for the unit.
4. **Voltage control.** Controls the **rate** of heating, *not the temperature!* The higher the setting, the faster the temperature rise.
5. **Stage light.** Provides illumination for samples.
6. **Eyepiece.** Magnifies the sample.
7. **Thermometer.** Gives temperature of sample.
8. **Thermometer end cap.** Keeps thermometer from falling out. If the cap becomes loose, the thermometer tends to go belly-up, and the markings turn over. Don’t try to fix this while the unit is hot. Let it cool so you won’t get burned.
9. **The hot stage.** This is the heating block that samples are melted on.

**OPERATION OF THE FISHER-JOHNS APPARATUS**

1. *Don’t assume that the unit is cold.* That is a good way to get burned.
2. Keep your grubby fingers off the cover slides. Use tweezers or forceps.
3. Place a clean round glass cover slide in the well on the hot stage. *Never melt any samples directly on the metal stage.* Ever!

4. Put a few crystals on the glass. Not too many. As long as you can see them melt, you're all right.

5. Put another cover slide on top of the crystals to make a sandwich.

6. Set the voltage control to zero if it's not already there.

7. Turn on–off switch to ON. The light source should illuminate the sample. If not, call for help!

8. Now science turns into art. Set the voltage control to any convenient setting. The point is to get up to *within 20°C* of the *supposed* melting point. Yep, that's right. If you have no idea what the melting point is, it may require several runs as you keep skipping past the point with a temperature rise of 5–10°C per minute. A convenient setting is 40. This is just a suggestion, not an article of faith.

9. After you've melted a sample, let it cool, and remove the sandwich of sample and cover slides. *Throw it away!* Use an appropriate waste container.

10. Once you have an idea of the melting point (or looked it up in a handbook, or you were told), *get a fresh sample*, and bring the temperature up quickly at about 5–10°C per minute to *within 20°C* of this approximate melting point. Then turn down the voltage control to get a 2°C per minute rise. Patience!

11. When the first crystals *just start to melt*, record the temperature. When the last crystal *just disappears*, record the temperature. If both points appear to be the same, either the sample is extremely pure, or the temperature rise was *too fast*.

12. Turn the on–off switch to OFF. Now set the voltage control to zero.

13. Let the stage cool, then remove the sandwich.

**THE THOMAS–HOOVER APPARATUS**

The Thomas–Hoover apparatus (Fig. 36) is the electromechanical equivalent of the Thiele tube or open beaker and hot oil methods (see “Using the Thiele Tube”). It has lots of features, and you should look for the following.
**Fig. 36** The Thomas–Hoover apparatus.

1. **Light box.** At the top of the device, towards the back, a box holds a fluorescent light bulb behind the thermometer. On the right side of this box are the fluorescent light switches.

2. **Fluorescent light switches.** Two buttons. Press and hold the red button down for a bit to light the lamp; press the black button to turn the lamp off.
3. **Thermometer.** A special $300^\circ$ thermometer in a metal jacket is immersed in the oil bath that’s in the lower part of the apparatus. Two slots have been cut in the jacket to let light illuminate the thermometer scale from behind, and to let a thermometer periscope read the thermometer scale from the front.

4. **Thermometer periscope.** In front of the thermometer, this periscope lets you read a small magnified section of the thermometer scale. By turning the small knob at the lower right of this assembly, you track the movement of the mercury thread, and an image of the thread and temperature scale appear in a stationary mirror just above the sample viewing area.

5. **Sample viewing area.** A circular opening cut in the front of the metal case such that you can see your samples in their capillary tubes (and the thermometer bulb) all bathed in the oil bath. You put the tubes into the oil bath through the holes in the capillary tube stage.

6. **Capillary tube stage.** In a semicircle about the bottom of the jacketed thermometer, yet behind the thermometer periscope, are five holes through which you can put your melting point capillaries.

7. **Heat.** Controls the rate of heating, not the temperature. The higher the setting, the faster the temperature rise. At Hudson Valley Community College, we’ve had a stop put in and you can only turn the dial as far as the number 7. When it gets up to 10, you always smoke the oil. Don’t do that.

8. **Power on–off switch.** Turns the unit on or off.

9. **Stirrer control.** Sets the speed of the stirrer from low to high.

10. **Vibrator on–off switch.** Turns the vibrator on or off. It’s a spring-return switch so you must hold the switch in the on position. Let go, and it snaps off.

11. **Line cords.** One brings a.c. power to the heater, stirrer, sample light, and vibrator. The other cord brings power to the fluorescent light behind the thermometer. Be sure both cords are plugged into live wall sockets.

**OPERATION OF THE THOMAS–HOOVER APPARATUS**

1. If the fluorescent light for the thermometer is not lit, press the red button at the right side of the light box and hold it down for a bit to start the lamp. The lamp should remain lit after you release the button.
2. Look in the thermometer periscope, turn the small knob at the lower right of the periscope base, and adjust the periscope to find the top of the mercury thread in the thermometer. Read the temperature. Wait for the oil bath to cool if the temperature is fewer than 20 Celsius degrees below the approximate melting point of your compound. You'll have to wait for a room temperature reading if you have no idea what the melting point is. You don't want to plunge your sample into oil that is so hot it might melt too quickly, or at an incorrect temperature.

3. Turn the voltage control to zero if it isn't there already.

4. Turn the power on–off switch to ON. The oil bath should become illuminated.

5. Insert your capillary tube in one of the capillary tube openings in the capillary tube stage. This is not simple. Be careful. If you snap a tube at this point, the entire unit may have to be taken apart to remove the pieces. It appears you have to angle the tube toward the center opening and angle the tube toward you (as you face the instrument) at the same time (Fig. 37). It’s as if they were placed on the surface of a conical funnel.

6. Adjust the magnifying glass for the best view of your sample.
7. Turn the stirrer knob so that the mark on the knob is about half of the way between the SLOW and FAST markings on the front panel. That’s just a suggestion. I don’t have any compelling reasons for it.

8. Adjust the thermometer periscope to give you a good view of the top of the mercury thread in the thermometer.

9. Now science turns into art. Set the heat control to any convenient setting. The point is to get up to within 20°C of the supposed melting point. If you have no idea what the melting point is, it may require several runs as you keep skipping past the point with a temperature rise of 5–10°C per minute. A convenient setting is 4. This is just a suggestion, not an article of faith.

10. Remember, you’ll have to keep adjusting the thermometer periscope to keep the top of the mercury thread centered in the image.

11. After you’ve melted a sample, throw it away!

12. Once you have an idea of the melting point (or looked it up in a handbook, or were told), get a fresh sample, and bring the temperature up quickly at about 5–10°C per minute to within 20°C of this approximate melting point. Then turn down the heat control to get a 2°C per minute rise. Patience!

13. When the first crystals just start to melt, record the temperature. When the last crystal just disappears, record the temperature. If both points appear to be the same, either the sample is extremely pure, or the temperature rise was too fast. If you record the temperature with the horizontal index line in the mirror matched to the lines etched on both sides of the periscope window and the top of the mercury thread at the same time, you’ll be looking at the thermometer scale head on. This will give you the smallest error in reading the temperature (Fig. 38).

14. Don’t turn the control much past 7. You can get a bit beyond 250°C at that setting, and that should be plenty for any solid compound you might prepare in this lab. Above this setting, there’s a real danger of smoking the oil.

15. Turn the power switch to OFF. You can also set the heat control to zero for the next person.

16. Press the black button on the right side of the light box and turn the fluorescent light off.

17. Remove all capillary tubes.
There are a few more electric melting point apprati around, and much of them work the same. A **sample holder**, **magnifying eyepiece**, and **voltage control** are common, and an apparently essential feature of these devices is that dial markings are almost *never* temperature settings. That is, a setting of 60 will not give a temperature of 60°C, but probably much higher.

**USING THE THIELE TUBE**

With the Thiele tube (Fig. 39) you use hot oil to transfer heat evenly to your sample in a melting point capillary, just like the metal block of the Mel-Temp apparatus does. You heat the oil in the sidearm and it expands. The hot oil goes up the sidearm, warming your sample and thermometer as it touches them. Now, the oil is cooler and it falls to the bottom of the tube where it is heated again by a burner. This cycle goes on automatically as you do the melting point experiment in the Thiele tube.
Oil cools, falls to bottom and recirculates.

**Notched cork holds thermometer without pressure buildup**

**Thiele tube clamped here**

**Rubber ring above hot oil!**

**Heats sample in capillary tube**

**Hot oil rises**

**Heat here**

**Fig. 39** Taking melting points with the thiele tube.
Don’t get any water in the tube or when you heat the tube the water can boil and throw hot oil out at you. Let’s start from the beginning.

Cleaning the Tube

This is a bit tricky, so don’t do it unless your instructor says so. Also, check with your instructor before you put fresh oil in the tube.

1. Pour the old oil out into an appropriate container and let the tube drain.
2. Use a hydrocarbon solvent (hexane, ligroin, petroleum ether — and no flames!) to dissolve the oil that’s left.
3. Get out the old soap and water and elbow grease, clean the tube, and rinse it out really well.
4. Dry the tube in a drying oven (usually > 100°C) thoroughly. Carefully take it out of the oven and let it cool.
5. Let your instructor examine the tube. If you get the OK, then add some fresh oil. Watch it. First, no water. Second, don’t overfill the tube. Normally, the oil expands as you heat the tube. If you’ve overfilled the tube, oil will crawl out and get you.

Getting the Sample Ready

Here you use a loaded melting point capillary tube (see “Loading the Melting Point Tube”) and attach it directly to the thermometer. The thermometer, unfortunately, has bulges; there are some problems, and you may snap the tube while attaching it to the thermometer.

1. Get, or cut, a thin rubber ring from a piece of rubber tubing.
2. Put the bottom of the loaded M.P. tube just above the place where the thermometer constricts (Fig. 40), and carefully roll the rubber ring onto the M.P. tube.
3. Reposition the tube so that the sample is near the center of the bulb and the rubber ring is near the open end. Make sure the tube is vertical.

Dunking the Melting Point Tube

There are more ways of keeping the thermometer suspended in the oil than I care to list. You can cut or file a notch on the side of the cork, drill a hole, and
insert the thermometer (*Be careful!*) Finally, cap the Thiele tube (Fig. 39). The notch is there so that pressure will not build up as the tube is heated. *Keep the notch open, or the setup may explode.*

But this requires drilling or boring corks, something you try to avoid (why have ground glass jointware in the undergraduate lab?). You can *gently* hold a thermometer and a cork in a clamp (Fig. 41). Not too much pressure, though!

Finally, you might put the thermometer in the *thermometer adapter* and suspend that, clamped gently by the rubber part of the adapter, not by the ground glass end. Clamping ground glass will score the joint.

**Heating the Sample**

The appropriately clamped thermometer is set up in the Thiele tube as in (Fig. 39). Look at this figure *now* and remember to heat the tube *carefully* — *always carefully* — *at the elbow*. *Then:*

1. If you don’t know the melting point of the sample, heat the oil fairly quickly, *but no more than 10°C per minute* to get a rough melting point.
Fig. 41 Safely suspended thermometer with Thiele tube.

And it will be rough indeed, since the temperature of the thermometer usually lags that of the sample.

2. After this sample has melted, lift the thermometer and attached sample tube carefully (*it may be HOT*) by the thermometer up at the clamp, until they are *just out of the oil*. This way the thermometer and sample can cool, and the hot oil can drain off. Wait for the thermometer to cool to about room temperature before you remove it entirely from the tube. Wipe off some of the oil, reload a melting point tube (*never remelt melted samples*), and try again. And heat at 2°C per minute this time.
Recrystallization
The essence of a recrystallization is a **purification**. Messy, dirty, compounds are cleaned up, purified, and can then hold their heads up in public again. The sequence of events you use will depend a lot on how messy your crude product is, and just how soluble it will be in various solvents.

In any case, you’ll have to remember a few things.

1. Find a solvent that will **dissolve the solid while hot**.
2. The same solvent **should not dissolve it while cold**.
3. The **cold solvent** must keep impurities dissolved in it **forever or longer**.

This is the major problem. And it requires some experimentation. That’s right! Once again, art over science. Usually, you’ll know what you should have prepared, so the task is easier. It requires a trip to your **notebook**, and possible, a **handbook** (see Chapter 2, “Keeping a Notebook” and Chapter 3, “Interpreting a Handbook”). You have the data on the solubility of the compound in your notebook. What’s that you say? **You don’t have the data in your notebook?** Congratulations, you get the highest F in the course.

Information in the notebook (which came from a handbook) for your compound might say, for alcohol (meaning ethyl alcohol), **s.h.** Since this means soluble in hot alcohol, it implies insoluble in cold alcohol (and you wondered what the **i** meant). Then alcohol is probably a good solvent for recrystallization of that compound. Also, check on the **color** or **crystalline form**. This is important since

1. A color in a supposedly white product is an impurity.
2. A color in a colored product is **not** an impurity.
3. The wrong **color** in a product is an impurity.

You can usually assume impurities are present in small amounts. Then you don’t have to guess what possible impurities might be present or what they
might be soluble or insoluble in. If your sample is really dirty, the assumption can be fatal. This doesn’t usually happen in an undergraduate lab, but you should be aware of it.

**FINDING A GOOD SOLVENT**

If the solubility data for your compound are not in handbooks, then

1. Place 0.1 g of your solid (weighed to 0.01 g) in a test tube.
2. Add 3 ml of a solvent, stopper the tube, and shake the bejesus out of it. If all of the solid **dissolves at room temperature**, then your solid is **soluble**. Do **not** use this solvent as a recrystallization solvent. (You must make note of this in your notebook, though).
3. If none (or very little) of the solid dissolved at room temperature, unstopper the tube and heat it (Careful — no flames!) and shake it and heat it and shake it. You may have to heat the solvent to a gentle boil (Careful! Solvents with low boiling points often boil away). If it does **not** dissolve at all, then do not use this as a recrystallization solvent.
4. If the sample **dissolved when HOT**, and **did not dissolve at room temperature**, you’re on the trail of a good recrystallization solvent. One last test.
5. Place this last test tube in an ice-water bath, and cool it to about 5°C or so. If lots of crystals come out, this is good, and this is your recrystallization solvent.
6. Suppose your crystals don’t come back when you cool the solution. Get a glass rod into the test tube, stir the solution, rub the inside of the tube with the glass rod, agitate that solution. If crystals still don’t come back, perhaps you’d better find another solvent.
7. Suppose, after all this, you still haven’t found a solvent. Look again. Perhaps your compound **completely** dissolved in ethanol at room temperature, and would **not** dissolve in water. AHA! Ethanol and water are **miscible** (i.e., they mix in all proportions) as well. You will have to perform a **mixed-solvent recrystallization** (see “Working with a Mixed-Solvent System”).
GENERAL GUIDELINES FOR A RECRYSTALLIZATION

Here are some general rules to follow for purifying any solid compound.

1. Put the solid in an Erlenmeyer flask, not a beaker. If you recrystallize compounds in beakers, you may find the solid climbing the walls of the beaker to get at you as a reminder. A 125-ml Erlenmeyer usually works. Your solid should look comfortable in it, neither cramped, nor with too much space. You probably shouldn’t fill the flask more than one fifth to one fourth full.

2. Heat a large quantity of a proven solvent (see preceding) to the boiling point, and slowly add the hot solvent. Slowly! A word about solvents: Fire! Solvents burn! No flames! A hot plate here would be better. You can even heat solvents in a steam or water bath. But — No flames!

3. Carefully add the hot solvent to the solid to just dissolve it. This can be tricky, since hot solvents evaporate, cool down, and so on. Ask your instructor.

4. Add a slight excess of the hot solvent (5–10 ml) to keep the solid dissolved.

5. If the solution is only slightly colored, the impurities will stay in solution. Otherwise, the big gun, activated charcoal, may be needed (see “Activated Charcoal”). Remember, if you were working with a colored compound, it would be silly to try to get rid of all the color, since you would get rid of all the compound and probably all your grade.

6. Keep the solvent hot (not boiling) and look carefully to see if there is any trash in the sample. This could be old boiling stones, sand, floor sweepings, and so on. Nothing you’d want to bring home to meet the folks. Don’t confuse real trash with undissolved good product! If you add more hot solvent, good product will dissolve, and trash will not. If you have trash in the sample, do a gravity filtration (see following).

7. Let the Erlenmeyer flask and the hot solution cool. Slow cooling gives better crystals. Garbage doesn’t get trapped in them. But this can take what seems to be an interminable length of time. (I know, the entire lab seems to take an interminable length of time.) So, after the flask cools and it’s just warm to the touch, then put the flask in an ice-water bath to cool. Watch it! The flasks have a habit of turning over in the water baths.
and letting all sorts of water destroy all your hard work! Also, a really hot flask will shatter if plunged into the ice bath, so again, watch it.

8. When you’re through cooling, filter the crystals on a Buchner funnel.

9. Dry them and take a melting point, as described in Chapter 9.

**GRAVITY FILTRATION**

If you find yourself with a flask full of hot solvent, your product dissolved in it, along with all sorts of trash, this is for you. You’ll need more hot solvent, a ringstand with a ring attached, possibly a clay triangle, some filter paper, a clean, dry flask, and a stemless funnel. Here’s how gravity filtration works.

1. Fold up a filter cone out of a piece of filter paper (Fig. 42). It should fit nicely, within a single centimeter or so of the top of the funnel. For those who wish to filter with more panache, try using fluted filter paper (see “world famous fan-folded fluted filter paper,” Fig. 52).

2. Get yourself a stemless funnel, or, at least, a short-stemmed funnel. Why? Go ahead and use a stem funnel and watch the crystals come out in the stem as the solution cools, blocking up the funnel (Fig. 43).

3. Put the filter paper cone in the stemless funnel.

4. Support this in a ring attached to a ringstand (Fig. 44). If the funnel is too small and you think it could fall through the ring, you may be able to get a wire or clay triangle to support the funnel in the ring (Fig. 45).

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![Fig. 42](image.jpg) **Fig. 42** Folding filter paper for gravity filtration.
**Fig. 43** The too long a funnel stem—Oops!

**Fig. 44** The gravity filtration setup with a funnel that fits the iron ring.
5. Put the new, clean, dry flask under the funnel to catch the hot solution as it comes through. All set?

6. Get that flask with the solvent, product and trash hot again. (No flames!) You should get some fresh, clear solvent hot as well. (No flames!)

7. Carefully pour the hot solution into the funnel. As it is, some solvents evaporate so quickly that product will probably come out on the filter paper. It is often hard to tell the product from the insoluble trash. Then —

8. Wash the filter paper down with a little hot solvent. The product will redissolve. The trash won’t.

9. You now let the trash-free solution cool and clean crystals should come out. Since you have probably added solvent to the solution, don’t be surprised if no crystals come out of solution. Don’t panic either! Just boil away some of the solvent, let your solution cool, and wait for the crystals again. If they still don’t come back, just repeat the boiling.

**Do not boil to dryness!**

Somehow, lots of folk think recrystallization means dissolving the solid, then boiling away all the solvent to dryness. NO! There must be a way to convince these lost souls that the impurities will deposit on the crystals. After the solution has cooled, crystals come out, sit on the bottom of the flask, and must be covered by solvent! Enough solvent to keep those nasty impurities dissolved and off the crystals.
THE BUCHNER FUNNEL AND FILTER FLASK

The Buchner funnel (Fig. 46) is used primarily for separating crystals of product from the liquid above them. If you have been boiling your recrystallization solvents dry, you should be horsewhipped and forced to reread these sections on recrystallization!

1. Get a piece of filter paper large enough to cover all the holes in the bottom plate, yet not curl up the sides of the funnel. It is placed flat on the plate (Fig. 46).
2. Clamp a filter flask to a ringstand. This filter flask, often called a suction flask, is a very heavy-walled flask with a sidearm on the neck. A piece of heavy-walled tubing connects this flask to the water trap (see Fig. 48).

Fig. 46 The Buchner funnel at home and at work.
3. Now use a rubber stopper or filter adapter to stick the Buchner funnel into the top of the filter flask. The Buchner funnel makes the setup top-heavy and prone to be prone — and broken. Clamp the flask first, or go get a new Buchner funnel to replace the one you’ll otherwise break.

4. The water trap is in turn connected to a source of vacuum, most likely, a water aspirator (Fig. 47).

5. The faucet on the water aspirator should be turned on full blast! This should suck down the filter paper, which you now wet with some of the cold recrystallization solvent. This will make the paper stick to the plate. You may have to push down on the Buchner funnel a bit to get a good seal between the rubber adapter and the funnel.

6. Swirl and pour the crystals and solvent slowly, directly into the center of the filter paper, as if to build a small mound of product there. Slowly! Don’t flood the funnel by filling it right to the brim, and waiting for the level to go down. If you do that, the paper may float up, ruining the whole setup.

7. Use a very small amount of the same cold recrystallization solvent and a spatula to remove any crystals left in the flask. Then you can use some of the fresh, cold recrystallization solvent and slowly pour it over the crystals to wash away any old recrystallization solvent and dissolved impurities.

8. Leave the aspirator on and let air pass through the crystals to help them dry. You can put a thin rubber sheet, a rubber dam, over the funnel. The vacuum pulls it in and the crystals are pressed clean and dry. And you won’t have air or moisture blowing through, and possibly decomposing, your product. Rubber dams are neat.

9. When the crystals are dry, and you have a water trap, just turn off the water aspirator. Water won’t back up into your flask. [If you’ve been foolhardy and filtered without a water trap, just remove the rubber tube connected to the filter flask sidearm (Fig. 47)].

10. At this point, you may have a cake of crystals in your Buchner funnel. The easiest way to handle this is to carefully lift the cake of crystals out of the funnel along with the filter paper, plop the whole thing onto a larger piece of filter paper, and let the whole thing dry overnight. If you are pressed for time, scrape the damp filter cake from the filter paper, but
don't scrape any filter paper fibers into the crystals. Repeatedly press the crystals out between dry sheets of filter paper, changing sheets until the crystals no longer show any solvent spot after pressing. Those of you who use heat lamps may find your white crystalline product turning into instant charred remains.

11. When your cake is completely dried, weigh a vial, put in the product, and weigh the vial again. Subtracting the weight of the vial from the weight of the vial and sample will give the weight of the product. This weighing by difference is easier and less messy than weighing the crystals directly on the balance. This weight should be included in the label on your product vial (see Chapter 8, "On Products").

Just a Note

I've said that a Buchner funnel is used primarily for separating crystals of product from the liquid above them. And in the section on drying agents, I tell people to use a gravity filtration setup to separate a drying agent from a liquid product. Recently, I've had some people get the notion that you can Buchner filter products from drying agents. I don't advise that. You will probably lose a lot of your product, especially if it has a low boiling point (<100°C). Under this vacuum filtration your product simply evaporates along with your grade.

ACTIVATED CHARCOAL

Activated charcoal is ultrafinely divided carbon with lots of places to suck up big, huge, polar, colored impurity molecules. Unfortunately, if you use too much, it'll suck up your product! And, if your product was white, or yellow, it'll have a funny gray color from the excess charcoal. Sometimes, the impurities are untouched and only the product gets absorbed. Again, it's a matter of trial and error. Try not to use too much. Suppose you've got a hot solution of some solid, and the solution is highly colored. Well,

1. First, make sure your product should not be colored!
2. Take the flask with your filthy product off the heat and swirl the flask.
This dissipates any superheated areas so that when you add the activated charcoal, the solution doesn’t foam out of the flask and onto your shoes.

3. *Add the activated charcoal.* Put a small amount, about the size of a pea, on your spatula, then throw the charcoal in. Stir. The solution may turn black. Stir and heat.

4. Set up the *gravity filtration* and filter off the carbon. It is especially important to *wash off any product caught on the charcoal*, and it is really hard to see anything here. You should take advantage of *fluted filter paper.* It should give a more efficient filtration.

5. Yes, have some extra fresh solvent heated as well. You’ll need to add a few milliliters of this to the hot solution to help keep the crystals from coming out on the filter paper. And you’ll need more to help wash the crystals off of the paper when they come out on it anyway.

6. This solution should be *much cleaner* than the original solution. If not, you’ll have to *add charcoal and filter again.* There is a point of diminishing returns, however, and one or two treatments is usually all you should do. Get some guidance from your instructor.

Your solid products should not be gray. Liquid products (yes, you can do liquids!) will let you know that you didn’t get all the charcoal out. Often, you can’t see charcoal contamination in liquids while you’re working with them. The particles stay suspended for awhile, but after a few days, you can see a layer of charcoal on the bottom of the container. Sneaky, those liquids. By the time the instructor gets to grade all the products — *voila* — the charcoal has appeared.

**THE WATER ASPIRATOR: A VACUUM SOURCE**

Sometimes you’ll need a vacuum for special work like *vacuum distillation* and *vacuum filtration* as with the Buchner funnel. An inexpensive source of vacuum is the *water aspirator* (Fig. 47).

When you turn the water on, the water flow draws air in from the side port on the aspirator. The faster the water goes through, the faster the air is drawn in. Pretty neat, huh? I’ve shown a plastic aspirator, but many of the older metal varieties are still around.
You may have to pretest some aspirators before you find one that will work well. It'll depend upon the water pressure in the pipes, too. Even the number of people using aspirators on the same water line can affect the performance of these devices. You can test them by going to an aspirator and turning the faucet on full blast. It does help to have a sink under the aspirator. If water leaks out the side port, tell your instructor and find another aspirator. Wet your finger and place it over the hole in the side port to feel if there is any vacuum. If there is no vacuum, tell your instructor and find another aspirator. Some of these old, wheezing aspirators have a very weak vacuum. You must decide for yourself if the suction is "strong enough." There should be a splash guard or rubber tubing leading the water stream directly into the sink. This will keep the water from going all over the room. If you check and don't find such protection, see your instructor. All you have to do with a fully tested and satisfactory aspirator is hook it up to the water trap.
THE WATER TRAP

Every year I run a chem lab and when someone is doing a vacuum filtration, suddenly I’ll hear a scream and a moan of anguish, as water backs up into someone’s filtration system. Usually there’s not much damage, since the filtrate in the suction flask is generally thrown out. For vacuum distillations, however, this suck-back is disaster. It happens whenever there’s a pressure drop on the water line big enough to cause the flow to decrease so that there is a greater vacuum in the system than in the aspirator. Water, being water, flows into the system. Disaster.

So, for your own protection, make up a water trap from some stoppers, rubber tubing, a thick-walled Erlenmeyer or filter flask, and a screw clamp (Fig. 48). Do not use garden variety Erlenmeyers; they may implode without warning. Two versions are shown. I think the setup using the filter flask is more flexible. The screw clamp allows you to let air into your setup at a controlled rate. You might clamp the water trap to a ringstand when you use it. The connecting hoses have been known to flip unsecured flasks, two out of three times.

WORKING WITH A MIXED-SOLVENT SYSTEM—THE GOOD PART

If, after sufficient agony, you cannot find a single solvent to recrystallize your product from, you may just give up and try a mixed-solvent system. Yes, it does mean you mix more than one solvent, and recrystallize using the mixture. It should only be so easy. Sometimes, you are told what the mixture is and the correct proportions. Then it is easy.

For an example, I could use “solvent 1” and “solvent 2,” but that’s clumsy. So I’ll use a rethe ethanol–water system and point out the interesting stuff as I go along.

The Ethanol–Water System

If you look up the solubility of water in ethanol (or ethanol in water) you find an $\infty$. This means they mix in all proportions. Any amount of one dissolves
completely in the other—no matter what. Any volumes, any weights. You name it. The special word for this property is **miscibility**. Miscible solvent systems are the kinds you should use a mixed solvents. They keep you out of trouble. You'll be adding amounts of water to the ethanol, and ethanol to the water. If the two weren't miscible, they might begin to separate and form two layers as you changed the proportions. Then you'd have REAL trouble. So, go ahead. You can work with mixtures of solvents that aren't miscible. But don't say you haven't been warned.

The ethanol—water mixture is so useful because

1. **At high temperatures, it behaves like alcohol!**
2. **At low temperatures, it behaves like water!**
From this, you should get the idea that it would be good to use a mixed solvent to recrystallize compounds that are soluble in alcohol yet insoluble in water. You see, each solvent alone cannot be used. If the material is soluble in the alcohol, not many crystals come back from alcohol alone. If the material is insoluble in water, you cannot even begin to dissolve it. So, you have a mixed solvent, with the best properties of both solvents. To actually perform a mixed-solvent recrystallization you

1. Dissolve the compound in the smallest amount of hot ethanol.
2. Add hot water until the solution turns cloudy. This cloudiness is tiny crystals of compound coming out of solution. Heat this solution to dissolve the crystals. If they do not dissolve completely, add a very little hot ethanol to force them back into solution.
3. Cool, and collect the crystals on a Buchner funnel.

Any solvent pair that behaves the same way can be used. The addition of hot solvents to one another can be tricky. It can be extremely dangerous if the boiling points of the solvents are very different. For the water–methanol mixed solvent, if 95°C water hits hot methanol (B.P. 65.0°C), watch out!

There are other miscible, mixed-solvent pairs, pet. ether and diehyl ether, methanol and water, and ligroin and diethyl ether among them.

A MIXED-SOLVENT SYSTEM—THE BAD PART

Every silver lining has a cloud. More often than not, compounds “recrystallized” from a mixed-solvent system don’t form crystals. Your compound may form an oil instead.

Oiling out is what it’s called; more work is what it means. Compounds usually oil out if the boiling point of the recrystallization solvent is higher than the melting point of the compound, though that’s not the only time. In any case, if the oil solidifies, the impurities are trapped in the now solid “oil,” and you’ll have to purify the solid again.

Don’t think you won’t ever get oiling out if you stick to single, unmixed solvents. It’s just that with two solvents, there’s a greater chance you’ll hit upon a composition that will cause this.
Temporarily, you can

1. Add more solvent. If it’s a mixed-solvent system, try adding more of the solvent the solid is NOT soluble in. Or add more of the OTHER solvent. No contradiction. The point is to change the composition. Single solvent or mixed solvent, changing the composition is one way out of this mess.

2. Redissolve the oil by heating, then shake up the solution as it cools and begins to oil out. When these smaller droplets finally freeze out, they may form crystals that are relatively pure. They may not. You’ll probably have to clean them up again. Just don’t use the same recrystallization solvent.

Sometimes, once a solid oils out, it doesn’t want to solidify at all, and you might not have all day. Try removing a sample of the oil with an eyedropper or disposable pipette. Then get a glass surface (watch glass) and add a few drops of a solvent that the compound is known to be insoluble in (usually water). Then use the rounded end of a glass rod to triturate the oil with the solvent. Trituration can be described loosely as the beating of an oil into a crystalline solid. Then you can put these crystals back into the rest of the oil. Possibly they’ll act as seed crystals and get the rest of the oil to solidify. Again, you’ll still have to clean up your compound.

SALTING-OUT

Sometimes you’ll have to recrystallize your organic compound from water. No big deal. But sometimes your organic compound is more than ever so slightly insoluble in water, and not all the compound will come back. Solution? Salt solution! A pinch of salt in the water raises the ionic strength. There are now charged ions in the water. Some of the water that solvated your compound goes to be with the salt ions. Your organic compound does not particularly like charged ions anyway, so more of your organic compound comes out of the solution.

You can dissolve about 36 g of common salt in 100 ml of cold water. That’s
the upper limit for salt. You can estimate how much salt you’ll need to practically saturate the water with salt. Be careful though—if you use too much salt, you may find yourself collecting salt crystals along with your product (see also the application of salting-out when you have to do an extraction; “Extraction Hints”).

WORLD FAMOUS FAN-FOLDED FLUTED FILTER PAPER

Some training in origami is *de rigueur* for chemists. It seems that the regular filter paper fold is inefficient, since very little of the paper is exposed. The idea here is to **flute** or **corrugate** the paper, increasing the surface area in contact with your filtrate. You’ll have to do this several times to get good at it.

Right here let’s review the difference between **fold** and **crease**. Folding is folding; creasing is folding, then stomping on it, running fingers and fingernails over a fold over and over and over. Creasing so weakens the paper, especially near the point, that it may break at an inappropriate time in the filtration.

1. Fold the paper in half, then in half again, then in half again (Fig. 49). Press on this wedge of paper to get the fold lines to stay, but *don’t crease*. *Do this in one direction only.* Either always fold toward you or away from you, but not both.

![Fig. 49 Folding filter paper into eighths.](image-url)
2. Unfold this cone *twice* so it looks like a semicircle (Fig. 50), and put it down on a flat surface. Look at it and think for not less than two full minutes the first time you do this.

3. OK. Now try a “fan fold.” You alternately fold, first in one direction then the other, every individual eighth section of the semicircle (Fig. 51).

4. Open the fan and play with it until you get a fairly fluted filter cone (Fig. 52).

5. It’ll be a bit difficult, but try to find the two opposing sections that are NOT folded correctly. Fold them inward (Fig. 52), and you’ll have a fantastic fan-folded fluted filter paper of your very own.
Fold in *two opposite* corners

**Fig. 52** Finishing the final fluted fan.

P.S. For those with more money than patience, prefolded fan-folded fluted filter paper is available from suppliers.
Extraction is one of the more complex operations you’ll do in the organic chemistry lab. For this reason, I’ll go over it especially slowly and carefully. Another term you’ll see used simultaneously with extraction is washing. That’s because extraction and washing are really the same operation, but each leads to a different end. How else to put this?

Let’s make some soup. Put the vegetables, fresh from the store, in a pot. You run cold water in and over them to clean them and throw this water down the drain. Later, you run water in and over them to cook them. You keep this water— it’s the soup.

Both operations are similar. Vegetables in a pot in contact with water the first time is a wash. You remove unwanted dirt. You washed with water. The second time, vegetables in a pot in contact with water is an extraction. You’ve extracted essences of the vegetables into water. Very similar operations; very different ends.

To put it a little differently,

You would extract good material from an impure matrix.

You would wash the impurities from good material.

The vegetable soup preparation is a solid–liquid extraction. So is coffee making. You extract some component(s) of a solid directly into the solvent. You might do a solid–liquid extraction in lab as a separate experiment; liquid–liquid extractions are routine. They are so common that if you are told to do an extraction or a washing, it is assumed, you will use two liquids— two INSOLUBLE liquids— and a separatory funnel. The separatory funnel, called a sep funnel by those in the know, is a special funnel that you can separate liquids in. You might look at the section on separatory funnels (later in this chapter) right now, then come back later.

Two insoluble liquids in a separatory funnel will form layers; one liquid will float on top of the other. You usually have compounds dissolved in these layers, and either the compound you want is extracted from one to the other, or junk you don’t want is washed from one layer to the other.

Making the soup, you have no difficulty deciding what to keep or what to throw away. First you throw the water away; later you keep it. But this can change. In a sep funnel, the layer you want to keep one time may not be the layer you want to keep the next time. Yet, if you throw one layer away prematurely, you are doomed.
NEVER-EVER LAND

Never, never, never, never, ever throw away any layer, until you are absolutely sure you'll never need it again. Not very much of your product can be recovered from the sink trap!

I'm using a word processor, so I can copy this warning over and over again, but let's not get carried away. One more time, WAKE UP OUT THERE!

Never, never, never, never, ever throw away any layer, until you are absolutely sure you'll never need it again. Not very much of your product can be recovered from the sink trap!

STARTING AN EXTRACTION

To do any extraction, you'll need two liquids, or solutions. They must be insoluble in each other. Insoluble here has a practical definition:

When mixed together, the two liquids form two layers.

One liquid will float on top of the other. A good example is ether and water. Handbooks say that ether is slightly soluble in water. When ether and water are mixed, yes, some of the ether dissolves; most of the ether just floats on top of the water.

Really soluble or miscible liquid pairs are no good for extraction and washing. When you mix them, they will not form two layers! In fact, they'll mix in all proportions. A good example of this is acetone and water. What kinds of problems can this cause? Well, for one, you cannot perform any extraction with two liquids that are miscible.

Let's try it. A mixture of say, some mineral acid (is HCl all right?) and an organic liquid, “Compound A,” needs to have that acid washed out of it. You dissolve the compound A–acid mixture in some acetone. It goes into the sep funnel, and you now add water to wash out the acid.

Acetone is miscible in water. No layers form! You lose!
Back to the lab bench. Empty the funnel. Start over. This time, having called yourself several colorful names because you should have read this section thoroughly in the first place, you dissolve the Compound A–acid mixture in ether and put it into the sep funnel. Add water, and two layers form! Now you can wash the acid from the organic layer to the water layer. The water layer can be thrown away.

Note that the acid went into the water, then the water was thrown out! So we call this a wash. If the water layer had been saved, we’d say the acid had been extracted into the water layer. It may not make sense, but that’s how it is.

Review:

1. You must have two insoluble liquid layers to perform an extraction.
2. Solids must be dissolved in a solvent, and that solvent must be insoluble in the other extracting or washing liquid.
3. If you are washing or extracting an organic liquid, dissolve it into another liquid, just like a solid, before extracting or washing it.

So these terms, extraction and washing are related. Here are a few examples.

1. Extract with ether. Throw ether together with the solution of product and pull out only the product into the ether.
2. Wash with 10% NaOH. Throw 10% NaOH together with the solution of product and pull out everything but product into the NaOH.
3. You can even extract with 10% NaOH.
4. You can even wash with ether.

So extraction is pulling out what you want from all else! Washing is pulling out all else from what you want.

And please note — you ALWAYS do the pulling from ONE LAYER INTO ANOTHER. That’s also two immiscible liquids.

You’ll have to actually do a few of these things before you get the hang of it, but bear with me. When your head stops hurting, reread this section.
Just before I go on to the separatory funnel, I'd like to comment on a few questions I keep hearing when people do washings and extractions.

1. **“Which layer is the water layer?”** Look at both layers in the funnel and get an idea of how big they are in relation to one another. Now add water to the funnel. Watch where the water goes. Watch which layer grows. Water to water. That's how you find the water (aqueous) layer. Don't rely on odor or color. Enough ether dissolves in water to give the water layer the odor of an ether layer; just enough of a highly colored substance in the wrong layer can mislead you.

2. **“How come I got three layers?”** Sometimes, when you pour fresh water or some other solvent into the funnel, you get a small amount hanging at the top, and it looks like there are three different layers. Yes, it looks as if there are three different layers, but there are not three different layers. Only two layers, where part of one has lost its way. Usually, this mysterious third layer looks just like its parent, and you might gently swirl the funnel and its contents to reunite the family.

3. **“What's the density of sodium hydroxide?”** You've just done a wash with 5–10% sodium hydroxide solution, you've just read something about finding various layers in the funnel by their densities, and, by this question, you've just shown that you've missed the point. Most wash solutions are 5 to 10% active ingredient dissolved in water. This means they are 90 to 95% water. Looking up the density of the solid reagents then, is a waste of time. The density of these solutions is very close to that of water. (10% NaOH has a specific gravity of 1.1089.)

4. **“I've washed this organic compound six times with sodium bicarbonate solution so why's it not basic yet?”** This involves finding the pH of the organic layer. I'll give it away right now. You cannot find the pH of an organic layer. Not directly. You find the pH of the aqueous layer that's been in contact with the organic layer. If the aqueous layer is on the top, dip a glass rod into it and touch the glass rod to your test paper. If the aqueous layer is on the bottom and your sev funnel in a ring, let a drop of the aqueous layer out of the funnel to hang on the outlet tip. Transfer the drop to your test paper. Warning. Be sure you are testing the
aqueous layer. Some organics are very tenacious and can get onto your glass rod. The organic layer may WET the test paper, but without water any color you see doesn’t mean much.

THE SEPARATORY FUNNEL

Before going on to some practical examples, you might want to know more about where all this washing and extracting is carried out. I’ve mentioned that it’s a special funnel called a separatory funnel (Fig. 53) and that you can impress your friends by calling it a sep funnel. Here are a few things you should know.

The Stopper

At the top of the sep funnel is a T glass stopper. There is a number, commonly T 22, possibly T 19/22, on the stopper head. Make sure this number is on the head and that it is the same as the number marked on the funnel. If this stopper is not so marked, you may find the product leaking over your shoes when you turn the sep funnel upside down. Try not to grease this stopper unless you plan to sauté your product. Unfortunately, these stoppers tend to get stuck in the funnel. The way out is to be sure you don’t get the ground glass surfaces wet with product. How? Pour solutions into the sep funnel as carefully as you might empty a shotglass of Scotch into the soda. Maybe use a funnel. To confuse matters, I’ll suggest you use a light coating of grease. Unfortunately, my idea of light and your idea of light may be different.

Consult your instructor!

The Glass Stopcock

This is the time-honored favorite of separatory funnel makers everywhere. There is a notch at the small end that contains either a rubber ring or a metal clip, but not both! There are two purposes for the ring.
1. To keep the stopcock from falling out entirely. Unfortunately, the rubber rings are not aware of this and the stopcock falls out anyway.

2. To provide a sideways pressure, pulling the stopcock in, so that it will not leak. Names and addresses of individuals whose stopcocks could not possibly leak and did so anyway will be provided on request. So provide a little sideways pressure of your own.

When you grease a glass stopcock (and you must), do it very carefully so
that the film of grease does not spread into the area of the stopper that comes in contact with any of your compound (Fig. 54).

The Teflon Stopcock

In wide use today, the Teflon stopcock (Fig. 55) requires no grease and will not freeze up! The glass surrounding the stopcock is not ground glass and cannot be used in funnels that require ground glass stopcocks! The Teflon stopcocks are infinitely easier to take care of. There is a Teflon washer, a rubber ring, and, finally, a Teflon nut, placed on the threads of the stopcock. This nut holds the whole thing on. Any leakage at this stopcock results from

1. A loose Teflon nut. Tighten it.
2. A missing Teflon washer or rubber ring. Have it replaced.

Fig. 54 The infamous glass stopcock.
3. An attempt to place the wrong size or taper Teflon stopcock into the funnel. This is extremely rare. Get a new funnel.

**Emergency stopcock warning!**

Teflon may not stick, *but it sure can flow!* If the stopcock is extremely tight, the Teflon will bond itself to all the nooks and crannies in the glass in interesting ways. When you’re through, always loosen the Teflon nut and “pop the stopcock” by pulling on the handle. The stopcock should be loose enough to spin freely when spun with one finger — *then remember to tighten it again before you use it.*

It seems to me that I’m the only one that reads the little plastic bags that hold the stopcock parts. Right on the bags it shows that after the stopcock goes in, *the Teflon washer goes on the stem first,* followed by the rubber ring, and then the Teflon nut (Fig. 55). So why do I find most of these things put together incorrectly?

**THE STEM**

The stem on a sep funnel can either be straight or have a ground glass joint on the end (Fig. 53). The ground glass joint fits the other jointware you may have and can be used that way as an addition funnel to add liquids or solutions
into a setup (see "Addition and Reflux"). You can use this type of separatory funnel as a sep funnel. You can't, however, use the straight-stem separatory funnel as an addition funnel without some help; remember, straight glass tubes don't fit ground glass joints (see "The Adapter With Lots of Names").

WASHING AND EXTRACTING VARIOUS THINGS

Now, getting back to extractions, there are really only four classes of compounds that are commonly handled in undergraduate extractions or washings.

1. **Strong Acids.** The mineral acids, and organic acids (e.g., benzoic acid). You usually extract these into sodium bicarbonate solution or wash them with it.

2. **Really weak acids.** Usually phenols, or substituted phenols. Here, you'd use a sodium hydroxide solution for washing or extraction. You need a strong base to work with these weak acids.

3. **Organic bases.** Any organic amine (aniline, triethylamine, etc.). As you use bases to work with acids, use a dilute acid (5 to 10% HCl, say) to extract or wash these bases.

4. **Neutral compounds.** All else, by these definitions (e.g., amides, ethers, alcohols, hydrocarbons).

HOW TO EXTRACT AND WASH WHAT

Here are some practical examples of washings and extractions, covering various types and mixtures and separations and broken down into the four classifications listed above.

1. **A strong organic acid.** Extract into sat'd (saturated) sodium bicarbonate solution.
(CAUTION! Foaming and fizzing and spitting and all sorts of carrying on.) The weak base turns the strong acid into a salt, and the salt dissolves in the water–bicarbonate solution. Because of all the fizzing, you’ll have to be very careful. Pressure can build up and blow the stopper out of the funnel. Invert the funnel. Point the stem AWAY FROM EVERYONE up and toward the BACK OF THE HOOD — and open the stopcock to vent or “burp” the funnel.

a. To recover the acid, add conc. (concentrated) HCl until the solution is acidic. Use pH or litmus paper to make sure. Yes, the solution really fizzes and bubbles. You should use a large beaker so material isn’t thrown onto the floor if there’s too much foam.

b. To wash out the strong acid, just throw the solution of bicarbonate away.

2. A weakly acidic organic acid. Extract into 10% NaOH–water solution. The strong base is needed to rip the protons out of weak acids (they don’t want to give them up) and turn them into salts; Then they’ll go into the NaOH–water layer.

a. To recover the acid, add conc. HCl until the solution of base is acid when tested with pH or litmus paper.

b. To wash out the weak acid, just throw this NaOH–water solution away.

3. An organic base. Extract with 10% HCl–water solution. The strong acid turns the base into a salt (This turning the whatever into a salt that dissolves in the water solution should be pretty familiar to you by now. Think about it.). Then the salt goes into the water layer.

a. To recover the base, add ammonium hydroxide to the water solution until the solution is basic to pH or litmus paper. Note that this is the reverse of the treatment given to organic acids.

b. To wash out an organic base, or any base, wash as above and throw out the solution.
4. **A neutral organic.** If you’ve extracted strong acids first, then weak acids, then bases, there are only neutral compound(s) left. If possible, just remove the solvent that now contains *only* your neutral compound. If you have *more than one neutral compound*, you may want to extract one from the other(s). You’ll have to find *two different immiscible organic liquids*, and *one liquid must dissolve ONLY the neutral organic compound you want!* A tall order. You must count on one neutral organic compound being *more soluble in one layer than in the other*. Usually the separation is *not clean — not complete*. And you have to do more work.

What’s “more work”? That depends on the results of your extraction.

**The Road to Recovery — Back-Extraction**

I’ve mentioned *recovery* of the four types of extractable materials, but that’s not all the work you’ll have to do to get the compounds in shape for further use.

1. If the recovered material is *soluble in the aqueous recovery solution*, you’ll have to do a **back-extraction**.

   a. Find a solvent that dissolves your compound, and *is not miscible in the aqueous recovery solution*. This solvent should boil at a low temperature (<100°C), since you will have to remove it. Ethyl ether is a common choice. *(Hazard! Very flammable).*

   b. *Then you extract your compound BACK FROM THE AQUEOUS RECOVERY SOLUTION into this organic solvent.*

   c. Dry this organic solution with a *drying agent* (see Chapter 7, “Drying Agents”).

   d. Now you can remove the organic solvent. Either distill the mixture or evaporate it, perhaps on a steam bath. All this is done away from flames and in a hood.

When you’re through removing the solvent and your product is not pure, clean it up. If your product is a liquid, you might distill it; if a solid, you might recrystallize it. Make sure it is clean.
2. If the recovered material is *insoluble in the aqueous recovery solution*, and it is a solid, collect the crystals on a Buchner funnel. If they are *not pure*, you should recrystallize them.

3. If the recovered material is *insoluble in the aqueous recovery solution* and it is a liquid, you can use your separatory funnel to *separate the aqueous recovery solution from your liquid product*. Then dry your liquid product and distill it if it is not clean. Or, you might just do a back-extraction as just described. This has the added advantage of getting out the small amount of liquid product that dissolves in the aqueous recovery solution and increases your yield. Remember to *dry* the back-extracted solution before you remove the organic solvent. Then distill your liquid compound if it is not clean.

A SAMPLE EXTRACTION

I think the only way I can bring this out is to use a typical example. This may ruin a few lab quizzes, but if it helps, it helps.

Say you have to separate a mixture of *benzoic acid* (1), *phenol* (2), *p-toluidine* (4-Methylanaline) (3), and *anisole* (methoxybenzene) (4) by extraction. The numbers refer to the class of compound, as previously listed. We’re assuming that none of the compounds react with any of the others and that you know we’re using all four types as indicated. Phenol and 4-methylanaline are corrosive toxic poisons and if you get near these compounds in lab, *be very careful*. When they are used as an example on these pages, however, you are quite safe. Here’s a sequence of tactics.

1. Dissolve the mixture in ether. Ether is insoluble in the water solutions you will extract into. Ether happens to dissolve all four compounds. Aren’t you lucky? You bet! It takes lots of hard work to come up with the “typical student example.”

2. Extract the ether solution with 10% HCl. This converts only compound 3, the basic p-toluidine, into the hydrochloride salt, which dissolves in the 10% HCl layer. You have just *extracted a base with an acid solution*. Save this solution for later.

3. Now extract the ether solution with sat’d sodium bicarbonate solution.
Careful! Boy will this fizz! Remember to swirl the contents and release the pressure. The weak base converts only compound 1, the benzoic acid, to a salt, which dissolves in the sat'd bicarbonate solution. Save this for later.

4. Now extract the ether solution with the 10% NaOH solution. This converts the compound 2, weak acid, phenol, to a salt, which dissolves in the 10% NaOH layer. Save this for later. If you do this step before step 3, that is, extract with 10% NaOH solution before the sodium bicarbonate solution, both the weak acid, phenol, and the strong acid, benzoic acid, will be pulled out into the sodium hydroxide. Ha-Ha. This is the usual kicker they put in lab quizzes, and people always forget it.

5. The only thing left is the neutral organic compound dissolve in the ether. Just drain this into a flask.

So, now we have four flasks with four solutions with one component in each. They are separated. You may ask, “How do we get these back?”

1. The basic compound (3). Add ammonium hydroxide until the solution turns basic (test with litmus or pH paper). The p-toluidine, or organic base (3), is regenerated.

2. The strong acid or the weak acid (1,2). A bonus. Add dilute HCl until the solution turns acidic to an indicator paper. Do it to the other solution. Both acids will be regenerated.

3. The neutral compound (4). It’s in the ether. If you evaporate the ether (No flames!), the compound should come back.

Now, when you recover these compounds, sometimes they don’t come back in such good shape. You will have to do more work.

1. Addition of HCl to the benzoic acid extract will produce huge amounts of white crystals. Get out the Buchner funnel and have a field day! Collect all you want. But they won’t be in the best of shape. Recrystallize them. (Note: This compound is insoluble in the aqueous recovery solution.)

2. The phenol extract is a different matter. You see, phenol is soluble in water, and it doesn’t come back well at all. So, get some fresh ether,
extract the phenol from HCl solution to the ether and evaporate the ether. Sounds crazy, no? No. Remember, I called this a back-extraction and you’ll have to do this more often than you would like to believe. (Note: This compound is soluble in the aqueous recovery solution.)

3. The p-toluidine should return after the addition of ammonium hydroxide. Recrystallize it from ethanol so it also looks respectable again.

4. The neutral anisole happens to be a liquid (B.P. 155°C), and you’ll have to take care when you evaporate the ether so as not to lose much anisole. Of course, you shouldn’t expect to see any crystals. Now this neutral anisole liquid that comes back after you’ve evaporated the ether (no flames!) will probably be contaminated with a little bit of all of the other compounds that started out in the ether. You will have to purify this liquid, probably by a simple distillation.

You may or may not have to do all of this with the other solutions, or with any other solution you ever extract in your life. You must choose. Art over science. As confusing as this is, I have simplified it a lot. Usually you have to extract these solutions more than once, and the separation is not as clean as you’d like. Not 100%, but pretty good. If you are still confused, see your instructor.

PERFORMING AN EXTRACTION OR WASHING

1. Suspend a sep funnel in an iron ring.
2. Remove the stopper.
3. Make sure the stopcock is closed! You don’t really want to scrape your product off the benchtop.
4. Add the solution to be extracted or washed. Less than half full, please. Add the extraction or washing solvent. An equal volume is usually enough. The funnel is funnel shaped and the equal volumes won’t look equal.
5. Replace the stopper.
6. Remove the sep funnel from the iron ring. Hold stopper and stopcock
tightly. Pressure may build up during the next step and blow your product out onto the floor.

7. Invert the sep funnel (Fig. 56).

   **Point the stem up away from everyone — up into the back of a hood if at all possible!**

   Make *sure* the liquid has drained down away from the stopcock, then *slowly* open the stopcock. You may hear a woosh, possibly a pffft, as the pressure is released. This is due to the high vapor pressure of some solvents, or to a gas evolved from a reaction during the mixing. This can cause big trouble when you are told to neutralize acid, by washing with sodium carbonate or sodium bicarbonate solutions.

8. **CLOSE THE STOPCOCK!**

9. Shake the funnel gently, invert it, open the stopcock again.

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**Fig. 56** Holding a sep funnel so as not to get stuff all over.
10. Repeat the steps 8 and 9 until no more gas escapes.

11. If you see that you might get an emulsion — a fog of particles — with this gentle inversion, do NOT shake the funnel vigorously. You might have to continue the rocking and inverting motions 30 to 100 times, as needed, to get a separation. Check with your instructor and with the hints on breaking up emulsions (see “Extraction Hints,” following). Otherwise, shake the funnel vigorously about 10 times, to get good distribution of the solvents and solutes. Really shake it.

12. Put the sep funnel back in the iron ring.

13. Remove the glass stopper. Otherwise the funnel won’t drain and you’ll waste your time just standing there.

14. Open the stopcock and let the bottom layer drain off into a flask.

15. Close the stopcock, swirl the funnel gently, then wait to see if any more of the bottom layer forms. If so, collect it. If not, assume you got it all in the flask.

16. Let the remaining layer out into another flask.

To extract any layer again, return that layer to the sep funnel, add fresh extraction or washing solvent, and repeat this procedure starting from step 5.

Never, never, never, never, ever throw away any layer, until you are absolutely sure you’ll never need it again. Not very much of your product can be recovered from the sink trap!

EXTRACTION HINTS

1. Several smaller washings or extractions are better than one big one.

2. Extracting or washing a layer twice, perhaps thrice, is usually enough. Diminishing returns set in after that.

3. Sometimes you’ll have to find out which layer is the water layer. This is so simple, it confounds everyone. Add 2–4 drops of each layer to a test tube containing 1 ml of water. Shake the tube. If the stuff doesn’t dissolve in the water, it’s not an aqueous (water) layer. The stuff may sink to the
bottom, float on the top, do **both**, or even turn the water cloudy! It will **not**, however, dissolve.

4. **If only the top layer** is being extracted or washed, it **does not have to be removed from the funnel**, ever. Just drain off the bottom layer, then add more fresh extraction or washing solvent. Ask your instructor about this.

5. You can combine the extracts of a multiple extraction, if they have the **same material in them**.

6. **If you have to wash your organic compound with water, and the organic is slightly soluble in water**, try washing with **saturated salt solution**. The theory is that if all that salt dissolved in the water, what room is there for your organic product? This point is a favorite of quizmakers, and should be remembered. It’s the same thing that happens when you add salt to reduce the solubility of your compound during a crystallization (see “Salting-Out”).

7. **If you get an emulsion**, you have not two distinct layers, but a kind of a fog of particles. Sometimes you can break up the charge on the suspended droplets by adding a little salt, or some acid or base. Or add ethanol. Or add salt. Or stir the solutions slowly with a glass rod. Or gravity filter the entire contents of your separatory funnel through filter paper. Or laugh. Or cry. Emulsion-breaking is a bit of an art. Careful with the acids and bases though. They can react with your product and destroy it.

8. **If you decide to add salt to a sep funnel**, don’t add so much that it clogs up the stopcock! For the same reason, keep drying agents out of sep funnels.

9. Sometimes some material comes out, or will not dissolve in the two liquid layers, and hangs in there in the **interface**. It may be that there’s not enough liquid to dissolve this material. One cure is to **add more fresh solvent of one layer or the other**. The solid may dissolve. If there’s no room to add more, you may have to remove both (yes, both) layers from the funnel, and try to dissolve this solid in either of the solvents. It can be confusing. If the material does **not** redissolve, then it is a new compound and should be saved for analysis. You should see your instructor for that one.
And
Now —
Boiling
Stones
All you want to do is start a distillation. Instructor walks up and says,

"Use a boiling stone or it'll bump."
"But I'm only gonna..."
"Use a boiling stone or it'll bump."
"It's started already and..."
"Use a boiling stone or it'll bump."
"I'm not gonna go and..."

Suddenly—WOOSH! Product all over the bench! Instant failure. Next time you put a boiling stone in before you start. No bumping. But your instructor won't let you forget the time you did it your way.

Don't let this happen to you. Use a brand new boiling stone every time you have to boil a liquid. A close up comparison between a boiling stone and the inner walls of a typical glass vessel reveals thousands of tiny nucleating points on the stone where vaporization can take place, in contrast to the smooth glass surface that can hide unsightly hot spots and lead to BUMP-ING, a massive instantaneous vaporization that will throw your product all over.

CAUTION! Introduction of a boiling stone into hot liquid may result in instant vaporization and loss of product. Remove the heat source, swirl the liquid to remove hot spots, then add the boiling stone.

Used as directed, the boiling stone will relieve minor hot spots and prevent loss of product through bumping. So remember... whenever you boil, wherever you boil,

ALWAYS USE A FRESH BOILING STONE!

Don't be the last on your bench to get this miracle of modern science made exclusively from nature's most common elements.
Sources of Heat
Many times you’ll have to heat something. Don’t just reach for the Bunsen burner. That flame you start just may be your own. There are alternate sources you should think of first.

THE STEAM BATH

If one of the components boils below 70°C and you use a Bunsen burner, you may have a hard time putting out the fire. Use a steam bath!

1. Find a steam tap. It’s like a water tap, only this one dispenses steam (Caution! You can get burned.)
2. Connect tubing to the tap now. It’s going to get awfully hot in use. Make sure you’ve connected a piece that’ll be long enough to reach your steam bath.
3. Don’t connect this tube to the steam bath yet! Just put it into a sink. Because steam lines are usually full of water from condensed steam, drain the lines first, otherwise you’ll waterlog your steam bath.
4. Caution! Slowly open the steam tap. You’ll probably hear bonking and clanging as steam enters the line. Water will come out. It’ll get hotter and may start to spit.
5. Wait until the line is mostly clear of water, then turn off the steam tap. Wait for the tubing to cool.
6. Slowly, carefully, and cautiously, making sure the tube is not hot, connect the tube to the inlet of the steam bath. This is the uppermost connection on the steam bath.
7. Connect another tube to the outlet of the steam bath — the lower connection — and to a drain. Any water that condenses in the bath while you’re using it will drain out.

Usually, steam baths have concentric rings as covers. You can control the “size” of the bath by removing various rings.

Never do this after you’ve started the steam. You will get burned!
And don’t forget — round-bottom flasks should be about halfway in the bath. Whether you should let steam rise up all around the flask or not appears to be a matter of debate. Lots of steam will certainly steam up the lab and may expose you to corrosion inhibitors (morpholine) in the steam lines. You should not, however, have steam shooting out the sides of the bath, or any other place. (Fig. 57).

**THE BUNSEN BURNER**

The first time you get the urge to take out a Bunsen burner and light it up, *don’t*. You may blow yourself up. Please check with your instructor to see if you even need a burner. Once you find out that you *can* use a burner, assume that the person who used it last didn’t know much about burners, and take some precautions so as not to burn your eyebrows off.

Now Bunsen burners are not the only kind. There are *Tirrill burners* and *Meker burners* as well. Some are more fancy than others, but they work pretty much the same. So when I say *burner* anywhere in the text it could be any of them.

![Diagram of steam bath](image)

*Fig. 57* The steam bath in use.
1. Find the **needle valve**. This is at the base of the burner. Turn it fully clockwise (inward) to stop the flow of gas completely. If your burner doesn’t have a needle valve, it’s a traditional Bunsen burner and the gas flow has to be regulated at the bench stopcock (Fig. 58). This can be dangerous, especially if you have to reach over your apparatus and burner to turn off the gas. Try to get a different model.

2. There is a **moveable collar** at the base of the burner which controls air flow. For now, see that all the holes are closed (i.e., no air gets in).

3. Connect the burner to the bench stopcock by some tubing and turn the bench valve **full on**. The bench valve handle should be parallel to the outlet (Fig. 58).

4. Now, **slowly** open the needle valve. You may be just able to hear some gas escaping. Light the burner. *Mind your face!* Don’t look down at the burner as you open the valve.

5. You’ll get a wavy yellow flame, something you don’t really want. But at least it’ll light. Now open the air collar a little. The yellow disappears; a blue flame forms. This is what you want.

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**Fig. 58** More than you may care to know about burners.
6. Now, adjust the needle valve and collar (the adjustments play off each other) for a steady blue flame.

**Burner Hints**

1. Air does not burn. You must wait until the gas has pushed the air out of the connecting tubing. Otherwise, you might conclude that none of the burners in the lab work. Patience, please.

2. When you set up the distillation or reflux, don’t waste a lot of time raising and lowering the entire setup so the burner will fit. This is nonsense. Move the burner! Tilt it! (See Fig. 59). If you leave the burner motionless under the flask, you may scorch the compound and your precious product can become a “dark intractable material.”

*Fig. 59* Don’t raise the flask, lower the burner.
3. Placing a wire gauze between the flame and the flask spreads out the heat evenly. Even so, the burner may have to be moved around. Hot spots can cause star cracks to appear in the flask (see Chapter 4, "Round-Bottom Flasks").

4. *Never* place the flask in the ring without a screen (Fig. 60). The iron ring heats up faster than the flask and the flask cracks in the nicest line around it you’ve ever seen. The bottom falls off and the material is all over your shoes.

**THE HEATING MANTLE**

A very nice source of heat, the heating mantle takes some special equipment and finesse.

1. **Variable voltage transformer.** The transformer takes the quite lethal 120 V from the wall socket and can change it to an equally dangerous 0 to 120 V, depending on the setting on the dial. Unlike temperature settings on a Mel-Temp, on a transformer 0 means 0 V, 20 means 20 V, and so on. I like to start at 0 V and work my way up. Depending on how much heat you want, values from 40 to 70 seem to be good places to start.
Also, you'll need a cord that can plug into both the transformer and the heating mantle.

2. The traditional fiberglass heating mantle. An electric heater wrapped in fiberglass insulation and cloth that looks vaguely like a catcher's mitt (Fig. 61).

3. The Thermowell heating mantle. You can think of the Thermowell heating mantle as the fiberglass heating mantle in a can. In addition, there is a hard ceramic shell that your flask fits into (Fig. 62). Besides just being more mechanically sound, it'll help stop corrosive liquids from damaging the heating element if your flask cracks while you're heating it.

4. Things not to do

   a. Don't ever plug the mantle directly onto the wall socket! I know, the curved prongs on the mantle connection won't fit, but the straight prongs on the adapter cord will. Always use a variable voltage transformer and start with the transformer set to zero.

   b. Don't use too small a mantle. The only cure for this is to get one that fits properly. The poor contact between the mantle and the glass doesn't transfer heat readily and the mantle burns out.

   c. Don't use too large a mantle. The only good cure for this is to get one that fits properly. An acceptable fix is to fill the mantle with sand, after the flask is in, but before you turn the voltage on. Otherwise, the mantle will burn out.

HINT. When you set up a heating mantle to heat any flask, usually for distillation or reflux, put the mantle on an iron ring and keep it clamped a few inches above the desktop (Fig. 61). Then clamp the flask at the neck, in case you have to remove the heat quickly. You can just unscrew the lower clamp and drop the mantle and iron ring.

PROPORTIONAL HEATERS AND STEPLESS CONTROLLERS

In all these cases of heating liquids for distillation or reflux, we really control the electric power, not the heat or temperature directly. Power is applied to the
heating elements, they warm up, yet the final temperature is determined by the heat loss to the room, the air, and, most important, the flask you’re heating. There are several types of electric power controls

1. **The variable voltage transformer.** We’ve discussed this just previously. Let me briefly restate the case: Set the transformer to 50 on the 0 – 100 dial and you get 50% of the line voltage, all the time, night and day, rain or shine.

2. **The mechanical stepless controller.** This appears to be the inexpensive replacement for the variable voltage transformer. Inside one model there’s a small heating wire wound around a bimetal strip with a magnet at one end (Fig. 63). A plunger connected to the dial on the front panel changes the distance between the magnet and a metal plate. With a heating mantle attached, when you turn the device on, current goes through the small heating wire and the mantle. The mantle is now on full.
**Fig. 62** A Thermowell heating mantle.

**Fig. 63** Inside a mechanical stepless controller.
blast (120V out of 120V from the electric wall socket)! As the small heating wire warms the bimetal strip, the strip expands, distorts, and finally pulls the magnet from its metal plate, opening the circuit. The mantle now cools rapidly (0V out of 120V from the wall socket), along with the bimetal strip. Eventually, the strip cools enough to let the magnet get close to that metal plate, and—CLICK—everything’s on full tilt again.

The front panel control varies the duty cycle, the time the controller is full on, to the time the controller is full off. If the flask, contents, and heating mantle are substantial, it takes a long time for them to warm up and cool down. A setup like that would have a large thermal lag. With small setups (approx. 50 ml. or so), there is a small thermal lag and very wild temperature fluctuations can occur. Also, operating a heating mantle this way is just like repeatedly plugging and unplugging it directly into the wall socket. There are not many devices that easily take that kind of treatment.

3. The electronic stepless controller. Would you believe a light dimmer? The electronic controller has a triac, a semiconductor device, that lets fractions of the a.c. power cycle through to the heating mantle. The a.c. power varies like a sine wave, from 0 to 120V from one peak to the next. At a setting of 25%, the triac remains off during much of the a.c. cycle, finally turning on when the time is right (Fig. 64). Although the triac does turn “full-off and full-on,” it does so at times so carefully controlled, that the mantle never sees full line power (unless you deliberately set it there).
Light dimmer and heating mantle triac power control.

**Fig. 64**
Unfortunately, glass apparatus needs to be held in place with more than just spit and bailing wire. In fact you would do well to use clamps. Life would be simple if there were just one type of fastener, but that’s not the case.

1. **The simple buret clamp** (Fig. 65). Though popular in other chem labs, the simple buret clamp just doesn’t cut it for organic lab. The clamp is too short, and adjusting angles with the “locknut” (by loosening the locknut, swiveling the clamp jaws to the correct angle and tightening the locknut against the back stop, away from the jaws) is not a great deal of fun. If you’re not careful, the jaws will slip right around and all the chemicals in your flask will fall out.

2. **The simple extension clamp and clamp fastener** (Fig. 66). This two-piece beast is the second best clamp going. It is much longer (approx. 12 in.), so you can easily get to complex setups. By loosening the **clamp holder thumbscrew**, the clamp can be pulled out, or pushed back, or rotated to any angle. By loosening the **ringstand thumbscrew**, the clamp, along with the clamp holder, can move up and down.

3. **The three-fingered extension clamp** (Fig. 67). Truly the Cadillac of

![Fig. 65 The "barely adequate for organic lab" buret clamp.](image)
CLAMPS AND CLAMPING

Fig. 66 The extension clamp and clamp fastener.

Fig. 67 The three-fingered clamp with clamp fastener.
clamps with a price to match. They usually try to confuse you with *two thumbscrews* to tighten, unlike the regular extension clamp. This gives a bit more flexibility, at the cost of a slightly more complicated way of setting up. You can make life simple by opening the *two-prong bottom jaw* to a 10 to 20° angle *from the horizontal* and treating that jaw as fixed. This will save a lot of wear and tear when you set equipment up, but you can *always move the bottom jaw* if you have to.

**CLAMPING A DISTILLATION SETUP**

You’ll have to clamp many things in your life as a chemist, and one of the more frustrating setups to clamp is the *simple distillation* (see Chapter 15, “Distillation”). If you can set this up, you probably will be able to clamp other common setups without much trouble. Here’s how to go about setting up the simple distillation.

1. OK, get a *ringstand* and an *extension clamp* and *clamp fastener* and put them all together. What heat source? A Bunsen burner, and you’ll need more room than with a heating mantle (see Chapter 13, “Sources of Heat”). In any case, you don’t know where the receiving flask will show up; and then you might have to readjust the entire setup. Yes, you should have read the experiment before so you’d know about the heating mantles.

2. Clamp the flask (around the neck) a few inches up the ringstand (Fig. 68). We *are* using heating mantles and you’ll need the room underneath to drop the mantle in case it gets too hot. That’s why the flask is *clamped at the neck*. Yes. That’s where the flask is *ALWAYS clamped*, no matter what heat source, so it doesn’t fall when the mantle comes down. What holds the mantle? Extension ring and clamp fastener.

3. Remember, whether you set these up from left to right, or right to left — *distilling flask first*!

4. Add the 3-way adapter now (Fig. 69). Thermometer and thermometer adapter come later.

5. Now add the condenser. Get another ringstand, extension clamp, and clamp fastener. There. Estimate the angle and height the clamp will be at
when the setup is clamped. Try setting the two-pronged jaw at about a 30° angle to the extension rod and call that the fixed jaw. Now turn the clamp so that the two-prong “fixed” jaw is at the bottom. Now, this two-pronged jaw of the clamp acts as a cradle for the condenser. Since tightening the top jaw won’t move the bottom jaw, there won’t be too many problems.
6. Place the two-prong clamp jaw in line with the condenser. Attach the condenser to the three-way adapter (Fig. 70). Hold everything! Sure. OK, *loosen extension clamp holder thumbscrew*, turn clamp to correct angle, and tighten. Now the height — up just a bit — Good! The lower "fixed" jaw cradles the consenser. *Tighten the ringstand thumbscrew (Arrrgh!) Clamps tend to move up ever so slightly as you tighten the fastener on the ringstand.*

7. Bottom jaw supports condenser — check. Joint at the three-way adapter/condenser OK? Good. Tighten wing nut and *bring the single-prong jaw down* onto condenser (Fig. 70). Not too tight.
8. Back from the stockroom again. Having put the vacuum adapter on the end of the condenser, expecting it to stay there by magic, you’ll be more careful with the new one.


10. All the clamps set up, all the joints tight—now where is that thermometer adapter?
Fig. 71 Correctly clamping the vacuum adapter.
Distillation
This separation or purification of liquids by vaporization and condensation is a very important step in one of man’s oldest professions. The word “still” lives on as a tribute to the importance of organic chemistry. The important points are

1. **Vaporization.** Turning a liquid to a vapor.
2. **Condensation.** Turning a vapor to a liquid.

Remember these. They show up on quizzes.

But when do I use distillation? That is a very good question. Use the guidelines below to pick your special situation, and turn to that section. But you *should* read all the sections, anyway.

1. **Class 1: Simple distillation.** Separating liquids boiling BELOW 150°C at one atmosphere from
   a. nonvolatile impurities.
   b. another liquid boiling at least 25°C higher than the first. The liquids should dissolve in each another.

2. **Class 2: Vacuum Distillation.** Separating liquids boiling ABOVE 150°C at 1 atm from
   a. nonvolatile impurities.
   b. another liquid boiling at least 25°C higher than the first. They should dissolve in one another.

3. **Class 3: Fractional Distillation.** Separating liquid mixtures, soluble in each other, that boil at less than 25°C from each other at 1 atm.

4. **Class 4: Steam Distillation.** Isolating tars, oils, and other liquid compounds *insoluble*, or slightly soluble, *in water at all temperatures*. Usually natural products are steam distilled. They do *not* have to be liquids at room temperatures (*e.g.*, caffeine, a solid, can be isolated from green tea.).

Remember, these are guides. If your compound boils at 150.0001°C don’t scream that you MUST do a vacuum distillation or both you and your product will die. I expect you to have some judgment and to pay attention to your instructor’s specific directions.
DISTILLATION NOTES

1. *EXCEPT* for Class 4, steam distillation, two liquids that are to be separated must dissolve in each other. If they did not, they would form separable layers, which you could separate in a separatory funnel (see Chapter 11, “Extraction and Washing”).

2. Impurities can be either *soluble* or *insoluble*. For example, the material that gives cheap wine its unique bouquet is soluble in the alcohol. If you distill cheap wine, you get clear grain alcohol separated from the “impurities,” which are left behind in the distilling flask.

CLASS 1: SIMPLE DISTILLATION (Fig. 72)

For separation of liquids boiling below 150°C at 1 atm from

1. nonvolatile impurities.
2. another liquid boiling 25°C higher than the first liquid. *They must dissolve in each other.*

Sources of Heat

If one of the components boils below 70°C and you use a Bunsen burner, you may have a hard time putting out the fire. Use a steam bath or a heating mantle. Different distillations will require different handling (see Chapter 13, “Sources of Heat”). All the distillations always require heating, so the sources of heat chapter is really closely tied to this section. This goes for enlightenment on the use of boiling stones and clamps as well (see Chapter 12, “And Now — Boiling Stones” and Chapter 14, “Clamps and Clamping”).

The 3-Way Adapter

If there is any one place your setup will fall apart, here it is (Fig. 73). When you set up the jointware, it is important that you have all the joints line up. This is
tricky, since, as you push one joint together, another pops right out. If you’re not sure, call your instructor. Let him inspect your work. Remember,

**All joints must be tight!**

**The Distilling Flask**

Fill the distilling flask with the liquid you want to distill. You can remove the thermometer and thermometer adapter, fill the flask using a funnel, then put the thermometer and its adapter back in place.

If you’re doing a **fractional distillation** with a **column** (a class 3 distillation), you should’ve filled the flask **before** clamping the setup. (Don’t ever pour your mixture down a column. That’ll contaminate everything!) You’ll
just have to disassemble some of the setup, fill the flask, reassemble what you’ve taken down, and pray that you haven’t knocked all the other joints out of line.

*Don’t fill the distilling flask more than half full.* Put in a boiling stone if you haven’t already. These porous little rocks promote bubbling and keep the liquid from superheating and flying out of the flask. This flying around is called *bumping*. NEVER drop a boiling stone into hot liquid or you may be rewarded by having your body soaked in the hot liquid as it foams out at you.

*Make sure all the joints in your setup are tight.* Start the heat S-L-O-W-L-Y until gentle boiling begins and liquid starts to drop into the receiving flask at the rate of about 10 drops per minute. *This is important.* If nothing comes over, you’re not distilling, merely wasting time. You may have to turn up the heat to keep material coming over.

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*Fig. 73* The commonly camouflaged until it’s too late open joint.
The Thermometer Adapter

Read all about it. Ways of having fun with thermometer adapters have been detailed (see text accompanying Fig. 22).

The Ubiquitous Clamp

A word about clamps. *Use!* They can save you $68.25 in busted glassware (see Chapter 14, "Clamps and Clamping").

The Thermometer

Make sure the ENTIRE thermometer bulb is *below the sidearm of the 3-way adapter*. If you don’t have liquid droplets condensing on the thermometer bulb, the temperature you read is *nonsense*. Keep a record of the temperature of the liquid or liquids that are distilling. It’s a check on the purity. Liquid collected over a 2°C range is fairly pure. Note the similarity of this range with that of the *melting point* of a pure compound (see Chapter 9, "The Melting Point Experiment").

The Condenser

Always keep cold water running through the condenser, enough so that *at least the lower half is cold* to the touch. Remember that water should go *in the bottom* and *out of the top* (Fig. 72). Also, the water pressure in the lab may change from time to time and usually goes up at night, since little water is used then. So, if you are going to let condenser cooling water run overnight, tie the tubing on at the condenser and the water faucet with wire or something. And if you don’t want to flood out the lab, see that the outlet hose can’t flop out of the sink.

The Vacuum Adapter

It is important that the tubing connector remain *open to the air* or else the entire apparatus will, quite simply, explode.
WARNING: Do not just stick the vacuum adapter on the end of the condenser and hope that it will not fall off and break.

This is foolish. I have no sympathy for anyone who will not use clamps to save their own breakage fee. They deserve it.

The Receiving Flask

The receiving flask should be large enough to collect what you want. You may need several, and they may have to be changed during the distillation. Standard practice is to have ONE flask ready for what you are going to throw away and others ready to save the stuff that you want to save.

The Ice Bath

Why everyone insists on loading up a bucket with ice and trying to force a flask into this mess, I’ll never know. How much cooling do you think you’re going to get with just a few small areas of the flask barely touching ice? Get a suitable receptacle — a large beaker, enamelled pan, or whatever. It should not leak. Put it under the flask. Put some water in it. Now add ice. Stir. Serves four.

Ice bath really means ice-water bath

THE DISTILLATION EXAMPLE

Say you place 50 ml of liquid A (B.P. 50°C) and 50 ml of liquid B (B.P. 100°C) in 250 ml R.B. flask. You drop in a boiling stone, fit the flask in a distillation setup, and turn on the heat. Bubbling starts and soon droplets form on the thermometer bulb. The temperature shoots up from room temperature to about 35°C, and a liquid condenses and drips into the receiver. That’s bad. The temperature should be close to 50°C. This low-boiling material is the forerun of a distillation, and you won’t want to keep it.
Keep letting liquid come over until the temperature stabilizes at about 49°C. Quick! Change receiving flasks NOW!

The new receiving flask is on the condenser and the temperature is about 49°C. GOOD. Liquid comes over and you heat to get a rate of about 10 drops per minute collected in the receiver. As you distill, the temperature slowly increases to maybe 51°C then starts moving up rapidly.

Here you stop the distillation and change the receiver. Now in one receiver you have a pure liquid, B.P. 49–51°C. Note this boiling range. It is just as good a test of purity as a melting point is for solids (see Chapter 11, “The Melting Point Experiment”).

Always report a boiling point for liquids as routinely as you report melting points for solids. The boiling point is actually a boiling range, and should be reported as such:

“B.P. 49–51°C”

If you now put on a new receiver, and start heating again, you may discover more material coming over at 50°C! Find that strange? Not so. All it means is that you were distilling too rapidly and some of the low-boiling material was left behind. It is very difficult to avoid this situation. Sometimes it is best to ignore it, unless a yield is very important. You can combine this “new” 50°C fraction with the other good fraction.

For B, boiling at 100°C, merely substitute some different boiling points and go over the same story.

THE DISTILLATION MISTAKE

OK, you set all this stuff up to do a distillation. Everything’s going fine. Clamps in the right place. No arthritic joints, even the vacuum adapter is clamped on, and the thermometer is at the right height. There’s a bright golden haze on the meadow and everything’s going your way. So, you begin to boil the liquid. You even remembered the boiling stone. Boiling starts slowly, then more rapidly. You think, “This is it!” Read that temperature, now. Into the notebook: “The mixture started boiling at 26°C”

And you are dead wrong.
What happened? Just ask —

is there liquid condensing on the thermometer bulb??

NO!

So, congratulations, you’ve just recorded the room temperature. There are days when over half the class will report distillation temperatures as “Hey I see it start boiling now” temperatures. Don’t participate. Just keep watching as the liquid boils. Soon, droplets will condense on the thermometer bulb. The temperature will go up quickly, then stabilize. NOW read the temperature. That’s the boiling point.

CLASS 2: VACUUM DISTILLATION

For separation of liquids boiling above 150°C at 1 atm from

1. nonvolatile impurities.
2. another liquid boiling 25°C higher than the first liquid. They must dissolve in each other. This is like the simple distillation with the changes shown (Fig. 74).

Why vacuum distill? If the substances boil at high temperatures at 1 atm, they may decompose when heated. Putting a vacuum over the liquid makes the liquid boil at a lower temperature. With the pressure reduced, there are fewer molecules in the way of the liquid you are distilling. Since the molecules require less energy to leave the surface of the liquid, you can distill at a lower temperature, and your compound doesn’t decompose.

Pressure Measurement

If you want to measure the pressure in your vacuum distillation setup, you’ll need a closed-end manometer. There are a few different types, but they all work essentially the same way. I’ve chosen a “stick” type (Fig. 75). This particular model needs help from a short length of rubber tubing and a glass T to get connected to the vacuum distillation setup.
DISTILLATION

Fig. 74  A vacuum distillation set up.

1. Turn on the source of vacuum and wait a bit for the system to stabilize.
2. Turn the knob on the manometer so that the notch in the joint lines up with the inlet.
3. Wait for the mercury in the manometer to stop falling.
4. Read the difference between the inner and outer levels of mercury. This is the system pressure, literally in millimeters of mercury, that we now call torr.
5. Turn the knob on the manometer to disconnect it from the inlet. Don’t leave the manometer permanently connected. Vapors from your distillation, water vapor from the aspirator, and so on, may contaminate the mercury.

Manometer Hints

1. Mercury is toxic, the vapor from mercury is toxic, mercury spilled breaks into tiny globules that evaporate easily and are toxic, it’ll alloy with your
This difference is the pressure in the system (22mm - 10mm = 12 mm Hg or 12 torr)

**Fig. 75** A closed-end "stick" manometer.
jewelry, and so on. Be very careful not to expose yourself (or anyone else) to mercury.

2. If the mercury level in the inner tubes goes lower than that of the outer tube it does NOT mean that you have a negative vacuum. Some air or other vapor has gotten into the inner stick, and with the vacuum applied, the vapor expands and drives the mercury in the inner tube lower than that in the outer tube. This manometer is unreliable and you should seek a replacement.

3. If a rubber tube connected to the vacuum source and the system (or manometer) collapses, you’ve had it. The system is no longer connected to the vacuum source, and as air from the bleed tube or vapor from the liquid you’re distilling fills your distillation setup, the pressure in the system goes up. Occasionally test the vacuum hoses and if they collapse under vacuum, replace them with sturdier hoses that can take it.

Leaks

Suppose, by luck of the draw, you’ve had to prepare and purify 1-octanol (B.P. 195°C). You know if you simply distill 1-octanol, you run the risk of having it decompose, so you set up a vacuum distillation. You hook your setup to a water aspirator and water trap and attach a closed-end “stick” manometer. You turn the water for the aspirator on full-blast and open the stick manometer. After a few minutes, nothing seems to be happening. You pinch the tubing going to the vacuum distillation setup, (but not to the manometer) closing the setup off from the source of vacuum. Suddenly, the mercury in the manometer starts to drop. You release the tube going to the vacuum distillation setup, and the mercury jumps to the upper limit. You have air leaks in your vacuum distillation setup.

Air leaks can be difficult to find. At best, you push some of the joints together again and the system seals itself. At worst, you have to take apart all the joints and regrease every one. Sometimes you’ve forgotten to grease all the joints. Often a joint has been etched to the point that it cannot seal under vacuum, when it is perfectly fine for other applications. Please get help from your instructor.
Pressure and Temperature Corrections

You’ve found all the leaks and the pressure in your vacuum distillation setup is, say, 25 torr. Now you need to know the boiling point of your compound, 1-octanol, this time at 25 torr and not 760 torr. You realize it’ll boil at a lower temperature, but just how low? The handy nomographs in (Figs. 76 and 77) can help you estimate the new boiling point.

This time you have the boiling point at 760 torr (195°C) and the pressure you are working at (25 torr) so you

1. Find the boiling point at 760 (195°C) on line B (the middle one).
2. Find the pressure you’ll be working at (25 torr) on line C (the one on the far right). You’ll have to estimate this point.
3. Using a straightedge, line up these two points and see where the straightedge cuts the observed boiling point line (Line A, far left). I get about 95°C.

So a liquid that boils at 195°C at 760 torr will boil at about 95°C at 25 torr. Remember, this is an estimate.

Now suppose you looked up the boiling point of 1-octanol and all you found was: 98°. This means that the boiling point of 1-octanol is 98°C at 19 torr. Two things should strike you.

1. This is a higher boiling point at a lower pressure than we’d gotten from the nomograph.
2. I wasn’t kidding about this process being an estimation of the boiling point.

Now we have a case of having an observed boiling point at a pressure that is not 760 torr (1-octanol again; 98°C at 19 torr). We’d like to get to 25 torr, our working pressure. This requires a double conversion.

1. On the observed B.P. line (line A) find 98°C.
2. On the pressure in torr line (line C) find 19.
3. Using a straightedge, connect those points. Now read the B.P. corrected to 760 (line B): I get 210°C.
Estimate 25 torr (your working pressure)

1. Find the B.P. at 760 torr (195°C)
2. Connect these dots
3. B.P. corrected to 760 torr
4. Get your B.P. at 25 torr here! (approx. 95°C)

Observed B.P.

Fig. 76 (a) One point conversion.
Get your B.P. at 25 torr here

B.P. corrected to 760 torr

Pivot about the point on B to working pressure on C

Draw line connecting published pressure and temperature (98°C @ 19 torr)

Pressure in torr

Observed B.P.

(b) Two point conversion.
Now, using the 210°C point as a fulcrum, pivot the straightedge until the 210°C point on line B and the pressure you’re working at (25 torr) on line C line up. You see, you’re in the same position as in the previous example with a “corrected to 760 torr B.P.” and a working pressure.

5. See where the straightedge cuts the observed boiling point line (line A). I get 105°C.

So, we’ve estimated the boiling point to be about 105°C at 25 torr. The last time it was 95°C at 25 torr. Which is it? Better you should say you expect your
compound to come over from 95–105°C. Again, this is not an unreasonable expectation for a vacuum distillation.

The pressure–temperature nomograph is really just a simple, graphical application of the Clausius–Clapyron equation. If you know the heat of vaporization of a substance, and its normal boiling point, you can calculate the boiling point at another temperature. You do have to assume that the heat of vaporization is constant over the temperature range you’re working with, and that’s not always so. Where’s the heat of vaporization in the nomograph? There is one built in, built into the slopes and spacings on the paper. And, yes, that means that the heat of vaporization is forced to be the same for all compounds, be they alkanes, aldehydes, or ethers. So do not be surprised at the inaccuracies in this nomograph; be amazed that it works as well as it does.

Vacuum Distillation Notes

1. Read ALL the notes on class 1.
2. The thermometer can be replaced by a gas inlet tube. It has a long, fine capillary at one end (Fig. 74). This is to help stop the extremely bad bumping that goes along with vacuum distillations. The fine stream of bubbles through the liquid produces the same results as a boiling stone. Boiling stones are useless, since all the adsorbed air is whisked away by the vacuum and the nucleating cavities plug up with liquid. The fine capillary does not let in a lot of air, so we are doing a vacuum distillation anyway. Would you be happier if I called it a reduced pressure distillation? An inert gas (nitrogen?) may be let in if the compounds decompose in air.
3. If you can get a magnetic stirrer and magnetic stirring bar you won’t have to use the gas inlet tube approach. Put a magnetic stirring bar in the flask with the material you want to vacuum distill. Use a heating mantle to heat the flask and put the magnetic stirrer under the mantle. When you turn the stirrer on, a magnet in the stirrer spins, and the stirring bar (a Teflon-coated magnet) spins. Admittedly, stirring through a heating mantle is not easy, but it can be done. Stirring the liquid also stops the bumping.

Remember, first the stirring, then the vacuum, THEN the heat—or WOOSHI Got it?
4. Control of heating is extremely critical. I don’t know how to shout this loudly enough on paper. Always apply the vacuum first and watch the setup for awhile. Air dissolved or trapped in your sample or a highly volatile leftover (maybe ethyl ether from a previous extraction) can come flying out of the flask without the heat. If you heated such a setup a bit and then applied the vacuum, your sample would blow all over, possibly right into the receiving flask. Wait for the contents of the distilling flask to calm down before you start the distillation.

5. If you know you have low-boiling material in your compound, think about distilling it at atmospheric pressure first. If, say, half the liquid you want to vacuum distill is ethyl ether from an extraction, consider doing a simple distillation to get rid of the ether. Then the ether (or any other low-boiling compound) won’t be around to cause trouble during the vacuum distillation. If you distill first at 1 atm, let the flask cool before you apply the vacuum. Otherwise your compound will fly all over and probably will wind up, undistilled and impure, in your receiving flask.

6. Grease all joints, no matter what (see “Greasing the Joints”). Under vacuum, it is easy for any material to work its way into the joints and turn into concrete, and the joints will never, ever come apart again.

7. The vacuum adapter is connected to a vacuum source, either a vacuum pump or a water aspirator. Real live vacuum pumps are expensive and rare and not usually found in the undergraduate organic laboratory. If you can get to use one, that’s excellent. See your instructor for the details. The water aspirator is used lots, so read up on it.

8. During a vacuum distillation, it is not unusual to collect a pure compound over a 10–20°C temperature range. If you don’t believe it, you haven’t ever done a vacuum distillation. It has to do with pressure changes throughout the distillation because the setup is far from perfect. Although a vacuum distillation is not difficult, it requires peace of mind, large quantities of patience, and a soundproof room to scream in so as not to disturb others.

9. A Claisen adapter in the distilling flask allows temperature readings to be taken and can help stop your compound from splashing over into the distillation receiver (Fig. 78). Also, you could use a three-neck flask (Fig. 79). Think! And, of course, use some glassware too.
For separation of liquids, soluble in each other, that boil less than 25°C from each other, use fractional distillation. This is like simple distillation with the changes shown (Fig. 80).

**Fractional distillation** is used when the components to be separated boil within 25°C of each other. Each component is called a fraction. Clever where
they get the name, eh? This temperature difference is not gospel. And don’t expect terrific separations either. Let’s just leave it at close boiling points. How close? That’s hard to answer. Is an orange? That’s easier to answer. If the experiment tells you to “fractionally distill,” at least you’ll be able to set it up right.

**How this works**

If one distillation is good, two is better. And fifty? Better still. So you have lots and lots of little, tiny distillations occurring on the surfaces of the column packing, which can be glass beads, glass helices, ceramic pieces, metal chips, or even stainless-steel wool.

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**Fig. 79** Same multipurpose setup with a three-neck flask.
Fig. 80 The fractional distillation setup.
As you heat your mixture it boils, and the vapor that comes off this liquid is \textit{richer in the lower boiling component}. The vapor moves out of the flask and condenses, say, on the first centimeter of column packing. Now, the composition of the liquid still in the flask has changed a bit — it is \textit{richer in the higher boiling component}. As more of this liquid boils, more hot vapor comes up, mixes with the first fraction, and produces a new vapor of different composition — \textit{richer yet in the more volatile (lower-boiling) component}. And guess what? This new vapor condenses in the second centimeter of column packing. And again, and again, and again.

Now all these are \textit{equilibrium steps}. It takes some time for the fractions to move up the column, get comfortable with their surroundings, meet the neighbors . . . . And if you \textit{never} let any of the liquid–vapor mixture out of the column, a condition called \textit{total reflux}, you might get a single pure component at the top; namely, the lower-boiling, more volatile component all by itself! This is an ideal separation.

Fat lot of good that does you when you have to hand in a sample. So, you turn up the heat, let some of the vapor condense, and \textit{take off this top fraction}. This raises hell in the column. \textit{Nonequilibrium conditions abound} — mixing. \textit{Arrrrgh!} No more completely pure compound. And the faster you distill, the faster you let material come over, the higher your \textit{throughput} — the worse this gets. Soon you’re at \textit{total takeoff} and there is no time for an equilibrium to get established. And if you’re doing that, you shouldn’t even bother using a column.

You must strike a compromise. Fractionally distill as slowly as you can, keeping in mind that eventually the lab does end. Slow down your fractional distillations; I’ve found that 5–10 drops per minute coming over into the receiving flask is usually suggested. It will take a bit of practice before you can judge the best rate for the best separation. See your instructor for advice.

\textbf{Fractional Distillation Notes}

1. Read \textit{ALL} the notes on class 1.
2. Make \textit{sure} you have not confused the \textbf{column} with the \textbf{condenser}. The \textit{column is wider and has glass projections inside}, at the bottom, to hold up the packing.
3. \textit{Don’t break off the projections!}
4. Do not run water through the jacket of the column!

5. Sometimes, the column is used without the column packing. This is all right, too.

6. If it is necessary, and it usually is, push a wad of heavy metal wool down the column, close to the support projections, to support the packing chips. Sometimes the packing is entirely this stainless steel wool. You can see that it is self-supporting.

7. Add the column packing. Shake the column lightly to make sure none of the packing will fall out later into your distillation.

8. With all the surface area of the packing, a lot of liquid is held up on it. This phenomenon is called column holdup, since it refers to the material retained in the column. Make sure you have enough compound to start with, or it will all be lost on the packing.

9. A chaser solvent or pusher solvent is sometimes used to help blast your compound off the surface of the packing material. It should have a tremendously high boiling point relative to what you were fractionating. After you've collected most of one fraction, some of this material is left on the column. So, you throw this chaser solvent into the distillation flask, fire it up, and start to distill the chaser solvent. As the chaser solvent comes up the column, it heats the packing material, your compound is blasted off the column packing and more of your compound comes over. Stop collecting when the temperature starts to rise — that's the chaser solvent coming over now. As an example, you might expect p-xylene (B.P. 138.4°C) to be a really good chaser, or pusher, for compounds that boil less than, say, 100°C.

But you have to watch out for the deadly azeotropes.

**AZEOTROPE**

Once in a while, you throw together two liquids and find that you cannot separate part of them. And I don't mean because of poor equipment, or poor technique, or other poor excuses. You may have an azeotrope, a mixture with a constant boiling point.

One of the best known examples is the ethyl alcohol – water azeotrope. This
96% alcohol – 4% water solution will boil to dryness, at a constant temperature. It’s slightly scary, since you learn that a liquid is a pure compound if it boils at a constant temperature. And you thought you had it made.

There are two types of azeotrope. If the azeotrope boils off first, it’s a **minimum boiling azeotrope**. After it’s all gone, if there is any other component left, only then will that component distill.

If any of the components come off first, and then the azeotrope, you have a **maximum boiling azeotrope**.

**Quiz question:**

Fifty milliliters of a liquid boils at 74.8°C from the beginning of the distillation to the end. Since there is no wide boiling range, can we assume that the liquid is pure?

No. It may be a constant boiling mixture called an azeotrope.

You should be able to see that you have to be really careful in selecting those chaser, or pusher solvents mentioned. Sure, water (B.P. 100°C) is hot enough to chase ethyl alcohol (B.P. 78.3°C) from any column packing. Unfortunately, water and ethyl alcohol form an azeotrope and the technique won’t work. (*Please* see “Theory of Distillation,” Chapter 28.)

### CLASS 4: STEAM DISTILLATION

Mixtures of tars and oils must not dissolve well in water (well, not much, anyway), so we can steam distill them. The process is pretty close to simple distillation, but you should have a way of getting fresh hot water into the setup without stopping the distillation.

Why steam distill? If the stuff you’re going to distill is *only slightly soluble in water* and may decompose at its boiling point and the bumping will be terrible with a vacuum distillation, it is better to **steam distill**. Heating the compound in the presence of steam makes the compound boil at a lower temperature. This has to do with partial pressures of water and organic oils and such.

There are two ways of generating steam:
External Steam Distillation

In an external steam distillation, you lead steam from a steam line, through a water trap, and thus into the system. The steam usually comes from a steam tap on the benchtop. This is classic. This is complicated. This is dangerous.

1. Set up your external steam distillation apparatus in its entirety. Have everything ready to go. Have the substance you want to distill already in the distilling flask. This includes having the material you want to distill in the distilling flask, the steam trap already attached, condensers up and ready, a large receiving flask, and so on. All you should have to do is attach a single hose from the steam tap to your steam trap and start the steam.

2. Have your instructor check your setup before you start! I cannot shout this loudly enough on this sheet of paper. Interrupting an external steam distillation, just because you forgot your head this morning, is a real trial.

3. Connect a length of rubber tubing to your bench steam outlet and lead the rubber tubing into a drain.

4. Now, watch out! Slowly, carefully, open the steam stopcock. Often you’ll hear clanging, bonking, and thumping, and a mixture of rust, oil, and dirt-laden water will come spitting out. Then some steam bursts come out. Finally, a stream of steam. Congratulations. You have just bled the steam line. Now close the steam stopcock, wait for the rubber tubing to cool a bit, and then . . .

5. Carefully (Caution — may be HOT!) attach the rubber tubing from the steam stopcock to the inlet of your steam trap.

6. Open the steam trap drain, then carefully reopen the bench steam stopcock. Let any water drain out of the trap then carefully close the drain clamp. Be CAREFUL.

7. You now have steam going through your distillation setup, and as soon as product starts to come over, you’ll be doing an external steam distillation. Periodically open the steam trap drain (Caution — HOT!) and let the condensed steam out.

8. Apparently, you can distill as fast as you can let the steam into your setup, as long as all the steam condenses and doesn’t go out into the room.
Sometimes you need to hook two condensers together, making a very long supercondenser, when you steam distill. Check with your instructor.

9. When you’re finished (see “Steam Distillation Notes” following), turn off the steam, let the apparatus cool, and dismantle everything.

There are many types of steam traps you can use with your distillation setup. I’ve shown one (Fig. 81), but this is not the only one, and you may use something different. The point is to note the *steam inlet* and the *trap drain*, and how to use them.

**Internal Steam Distillation**

1. You can add hot water to the flask (Fig. 82) that will generate steam and thus provide an **internal source of steam**. This method is used almost exclusively in an undergraduate organic lab for the simple reason that it is so simple.

2. Add to the distilling flask at least 3 times as much water (maybe more) as sample. Do not fill the flask much more than half full (three quarters, maybe). You’ve got to be careful. Very careful.

3. Periodically add more *hot water* as needed. When the water boils and turns to steam, it also leaves the flask, carrying product.

**Steam Distillation Notes**

1. Read *ALL* the notes on class 1 distillations.

2. Collect some of the distillate, the stuff that comes over, in a small test tube. Examine the sample. If you see *two layers*, or *the solution is cloudy*, you’re not done. Your product is still coming over. Keep distilling and keep adding hot water to generate more steam. *If you don’t see any layers, don’t assume you’re done.* If the sample is slightly soluble in the water, the two layers or cloudiness might not show up. Try *salting-out*. This has been mentioned before in connection with extraction and recrystallization as well (see “Salting-Out” and “Extraction Hints”). Add some salt to the solution you’ve collected in the test tube, shake the tube to dissolve the salt, and if you’re lucky, more of your product may be
Fig. 81 One example of an external steam trap.
squeezed out of the aqueous layer, forming a *separate layer*. If that happens, *keep steam distilling* until the product does not come out when you treat a test solution with salt.

3. There should be two layers of liquid in the receiving flask at the end of the distillation. One is *mostly water*. The other is *mostly product*. To find out which is which, add a small quantity of water to the flask. The water will go into the water layer. (Makes sense.) Be very careful with this test, however; it is sometimes very hard to tell where the water has gone.

4. If you have to get more of your organic layer out of the water, you can do a *back-extraction* with an immiscible solvent (see "The Road to Recovery — Back-Extraction").
Just about 80% of the reactions in organic lab involve a step called refluxing. You use a reaction solvent to keep materials dissolved and at a constant temperature by boiling the solvent, condensing it, and returning it to the flask.

For example, say you have to heat a reaction to around 80°C for 17 hours. Well, you can stand there on your flat feet and watch the reaction all day. Me? I'm off to the reflux.

Usually, you'll be told what solvent to use, so selecting one should not be a problem. What happens more often is that you choose the reagents for your particular synthesis, put them into a solvent, and reflux the mixture. You boil the solvent and condense the solvent vapor so that ALL the solvent runs back into the reaction flask (see "Fractional Distillation"). The reflux temperature is near the boiling point of the solvent. To execute a reflux,

1. Place the reagents in a round-bottomed flask. The flask should be large enough to hold both the reagents and enough solvent to dissolve them, without being much more than half full.
2. You should now choose a solvent that
   a. Dissolves the reactants at the boiling temperature.
   b. Does not react with the reagents.
   c. Boils at a temperature that is high enough to cause the desired reaction to go at a rapid pace.
3. Dissolve the reactants in the solvent. Sometimes the solvent itself is a reactant. Then don't worry.
4. Place a condenser, upright, on the flask, connect the condenser to the water faucet, and run water through the condenser (Fig. 83). Remember—in at the bottom and out at the top.
5. Put a suitable heat source under the flask and adjust the heat so that the solvent condenses no higher than halfway up the condenser. You'll have to stick around and watch for a while, since this may take some time to get started. Once the reaction is stable, though, go do something else. You'll be ahead of the game for the rest of the lab.
6. Once this is going well, leave it alone until the reaction time is up. If it's an overnight reflux, wire the water hoses on so they don't blow off when you're not there.
7. When the reaction time is up, turn off the heat, let the setup cool, dismantle it, and collect and purify the product.

**A DRY REFLUX**

If you have to keep the atmospheric water vapor out of your reaction, you must use a **drying tube and the inlet adapter** in the reflux setup (Fig. 84). You can use these if you need to keep water vapor out of any system, not just the reflux setup.
Fig. 84  Reflux setup à la drying tube.
1. If necessary, clean and dry the drying tube. You don't have to do a thorough cleaning unless you suspect that the anhydrous drying agent is no longer anhydrous. If the stuff is caked inside the tube, it is probably dead. You should clean and recharge the tube at the beginning of the semester. Be sure to use anhydrous calcium chloride or sulfate. It should last one semester. If you are fortunate, indicating Drierite, a specially prepared anhydrous calcium sulfate, might be mixed in with the white Drierite. If the color is blue, the drying agent is good; if red, the drying agent is no longer dry, and you should get rid of it (see Chapter 7, “Drying Agents”).

2. Put in a loose plug of glass wool or cotton to keep the drying agent from falling into the reaction flask.

3. Assemble the apparatus as shown, with the drying tube and adapter on top of the condenser.

4. At this point, reagents may be added to the flask and heated with the apparatus. Usually, the apparatus is heated while empty to drive water off the walls of the apparatus.

5. Heat the apparatus, usually empty, on a steam bath, giving the entire setup a quarter-turn every so often to heat it evenly. A burner can be used if there is no danger of fire and if heating is done carefully. The heavy ground glass joints will crack if heated too much.

6. Let the apparatus cool to room temperature. As it cools, air is drawn through the drying tube before it hits the apparatus. The moisture in the air is trapped by the drying agent.

7. Quickly add the dry reagents or solvents to the reaction flask, and reassemble the system.

8. Carry out the reaction as usual like a standard reflux.

**ADDITION AND REFLUX**

Every so often you have to add a compound to a setup while the reaction is going on, usually along with a reflux. Well, you don't break open the system, let toxic fumes out, and make yourself sick to add new reagents. You use an addition funnel. Now, we talked about addition funnels back with separa-
tory funnels (Chapter 11) when we were considering the stem, and that might have been confusing.

Funnel Fun

Look at Fig. 83a. It is a true sep funnel. You put liquids in here and shake and extract them. But could you use this funnel to add material to a setup? NO. No ground glass joint on the end; and only glass joints fit glass joints. Right? Of course, right.

Figure 85c shows a pressure-equalizing addition funnel. See that sidearm? Remember when you were warned to remove the stopper of a separatory funnel so you wouldn’t build up a vacuum inside the funnel as you emptied it? Anyway, the sidearm equalizes the pressure on both sides of the liquid you’re adding to the flask, so it’ll flow freely, without vacuum buildup and without you having to remove the stopper. This equipment is very nice, very expensive, very limited, and very rare. And if you try an extraction in one of these, all the liquid will run out the tube onto the floor as you shake the funnel.

So a compromise was reached (Fig. 85b). Since you’ll probably do more extractions than additions, with or without reflux, the pressure-equalizing tube went out, but the ground glass joint stayed on. Extractions; no problem. The nature of the stem is unimportant. But during additions, you’ll have to take the responsibility to see that nasty vacuum buildup doesn’t occur. You can remove the stopper every so often or put a drying tube and inlet adapter in place of the stopper. The latter keeps moisture out and prevents vacuum buildup inside the funnel.

How to Set Up

There are at least two ways to set up an addition and reflux, using either a three-neck flask or a Claisen adapter. I thought I’d show both these setups with drying tubes. They keep the moisture in the air from getting into your reaction. If you don’t need them, do without them.

Often, the question comes up, “If I’m refluxing one chemical, how fast can I add the other reactant?” Try to follow your instructor’s suggestions. Anyway, usually the reaction times are fixed. So I’ll tell you what NOT, repeat NOT, to do.
If you reflux something, there should be a little ring of condensate, sort of a cloudy, wavy area in the barrel of the reflux condenser (Figs. 86 and 87). Assuming an exothermic reaction, the usual case, adding material from the funnel has the effect of heating up the flask. The ring of condensate begins to move up. Well, don’t ever let this get more than three quarters up the condenser barrel. If the reaction is that fast, a very little extra reagent or heating will push that ring out of the condenser and possibly into the room air. No No, no, no.
Fig. 86  Reflux and addition by Claisen tube.
ADDITION AND REFLUX

Fig. 87  Reflux and addition by three-neck flask
Sublimation
Sublimation occurs when you heat a solid and it turns directly into a vapor. It does not pass GO nor does it turn into a liquid. If you reverse the process — cool the vapor so that it turns back into a solid — you’ve condensed the vapor. Use the unique word, sublime, for the direct conversion of solid to vapor. Condense can refer to either vapor-to-solid or vapor-to-liquid conversions.

Figure 88 shows two forms of sublimation apparatus. Note all the similarities. Cold water goes in and down into a cold finger upon which the vapors

![Diagram of sublimation apparatus](image)

**Fig. 88** King-size and miniature sublimation apparatus.
from the crystals condense. The differences are that one is larger and has a ground glass joint. The sidearm test tube with cold-finger condenser is much smaller. To use them,

1. Put the crude solid into the bottom of the sublimator. How much crude solid? This is rather tricky. You certainly don’t want to start with so much that it touches the cold finger. And since as the purified solid condenses on the cold finger it begins to grow down to touch the crude solid, there has to be really quite a bit of room. I suggest that you see your instructor, who may want only a small amount purified.

2. Put the cold finger into the bottom of the sublimator. Don’t let the clean cold finger touch the crude solid. If you have the sublimator with the ground glass joint, lightly (and I mean lightly) grease the joint. Remember that greased glass joints should NOT be clear all the way down the joint.

3. Attach the hoses. Cold water goes in the center tube, pushing the warmer water out the side tube. Start the cooling water. Be careful!

4. If you’re going to pull a vacuum in the sublimator, do it now. If the vacuum source is a water aspirator, put a water trap between the aspirator and the sublimator. Otherwise you may get depressed if, during a sudden pressure drop, water backs up and fills your sublimator. Also, start the vacuum slowly. If not, air, entrained in your solid, comes rushing out and blows the crude product all over the sublimator, like popcorn.

5. When everything has settled down, slowly begin to heat the bottom of the sublimator, if necessary. You might see vapors coming off the solid. Eventually, you’ll see crystals of purified solid form on the cold finger. Since you’ll work with different substances, different methods of heating will have to be used. Ask your instructor.

6. Now the tricky parts. You’ve let the sublimator cool. If you’ve a vacuum in the sublimator, carefully — very carefully — introduce air into the device. A sudden inrush of air, and PLOP! Your purified crystals are just so much yesterday’s leftovers. Start again.

7. Now again, carefully — very carefully — remove the cold finger, with your pristine product clinging tenuously to the smooth glass surface, without a lot of bonking and shaking. Otherwise, PLOP! et cetera, et cetera, et cetera. Clean up and start again.
Chromatography: Some Generalities
Chromatography is perhaps the most useful means of separating compounds to purify and identify them. Indeed, separations of colored compounds on paper strips gave the technique its colorful name. Though there are many different types of chromatography, there are tremendously striking similarities among all the forms. Thin-layer, wet-column, and dry-column chromatography are common techniques you'll run across.

This chromatography works by differences in polarity. (That's not strictly true for all types of chromatography, but I don't have the inclination to do a 350-page dissertation on the subject, when all you might need to do is separate the differently colored inks in a black marker pen.)

** ADSORBANTS **

The first thing you need is an adsorbant, a porous material that can suck up liquids and solutions. Paper, silica gel, alumina (ultrafine aluminum oxide), corn starch, and kitty litter (unused) are all fine adsorbants. Only the first three are used for chromatography. You may or may not need a solid support with these. Paper hangs together, is fairly stiff, and can stand up by itself. Silica gel, alumina, corn starch, and kitty litter are more or less powders and will need a solid support to hold them.

Now you have an adsorbant on some support, or a self-supporting adsorbant, like a strip of paper. You also have a mixture of stuff you want to separate. So you dissolve the mixture in an easily evaporated solvent, like methylene chloride, and put some of it on the adsorbant. Zap! It is adsorbed! Stuck on and held to the adsorbant. But because you have a mixture of different things, and they are different, they will be held to the adsorbant in differing degrees.

** SEPARATION OR DEVELOPMENT **

Well, now there's this mixture, sitting on this adsorbant, looking at you. Now you start to run solvents through the adsorbant. Study the following list of solvents. Chromatographers call these solvents eluents.
THE ELUATROPIC SERIES

THE ELUATROPIC SERIES

Not at all like the World Series, the eluatropic series is simply a list of solvents arranged according to increasing polarity.

Some solvents arranged in order of increasing polarity

(Least polar) Pet. ether
Cyclohexane
Increasing Toluene
Chloroform
Polarity Acetone
Ethanol
(Most polar) Methanol

So you start running “pet. ether” (remember, petroleum ether, a mixture of hydrocarbons like gasoline — not a true ether at all). It’s not very polar. So it is not held strongly to the adsorbant.

Well, this solvent is traveling through the adsorbant, minding its own business, when it encounters the mixture placed there earlier. It tries to kick the mixture out of the way. But most of the mixture is more polar, held more strongly on the adsorbant. Since the pet. ether cannot kick out the compounds more polar than itself very well, most of the mixture is left right where you put it.

No separation.

Desperate, you try methanol, one of the most polar solvents. It is really held strongly to the adsorbant. So it comes along and kicks the living daylights out of just about all the molecules in the mixture. After all, the methyl alcohol is more polar, so it can move right along and displace the other molecules. And it does. So, when you evaporate the methanol and look, all the mixture has moved with the methanol, so you get one spot that moved, right with the solvent front.

No separation.
Taking a more reasonable stand, you try chloroform, because it has an intermediate polarity. The chloroform comes along, sees the mixture, and is able to push out, say, all but one of the components. As it travels, kicking the rest along, it gets tired and starts to leave some of the more polar components behind. After a while, only one component is left moving with the chloroform, and that may be dropped, too. So, at the end, there are several spots left, and each of them is in a different place from the start. Each spot is at least one different component of the entire mixture.

Separation. At last!

I picked these solvents for illustration. They are quite commonly used in this technique. I worry about the hazards of using chloroform, however, because it’s been implicated in certain cancers. Many other common solvents, too, are suspected to be carcinogens. In lab, you will either be told what solvent (eluent) to use or you will have to find out yourself, mostly by trial and error.
Thin-Layer Chromatography: TLC
Thin-layer chromatography (TLC) is used for identifying compounds and determining their purity. The most common adsorbant used is silica gel. Alumina is gaining popularity, with good reason. Compounds should separate the same on an alumina plate as on an alumina column, and column chromatography using alumina is still very popular. And, it is very easy to run test separations on TLC plates, rather than carrying out tests on chromatographic columns.

Nonetheless, both these adsorbants are powdered and require a solid support. Microscope slides are extremely convenient. To keep the powder from just falling off the slides, manufacturers add a gypsum binder (plaster). Adsorbants with the binder usually have a “G” stuck on the name or say “For thin-layer use” on the container.

Sometimes a fluorescent powder is put into the adsorbant to help with visualization later. The powder usually glows a bright green when you expose it to 254-nm wavelength ultraviolet (UV) light. You can probably figure out that if a container of silica gel is labeled Silica Gel G-254, you’ve got a TLC adsorbant with all the bells and whistles.

Briefly, you mix the adsorbant with water, spread the mix on the microscope slide in a thin layer, and let it dry, then activate the coating by heating the coated slide on a hot plate. Then you spot or place your unknown compound on the plate, let an eluent run through the adsorbant (development), and finally examine the plate (visualization).

## PREPARATION OF TLC PLATES

1. Clean and dry several microscope slides.
2. In an Erlenmeyer flask, weigh out some adsorbant, and add water.
   a. For silica gel use a 1:2 ratio of gel to water. About 2.5 g gel and 5 g water will do for a start.
   b. For alumina, use a 1:1 ratio of alumina to water. About 2.5 g alumina and 2.5 g water is a good start.
3. Stopper the flask and shake it until all the powder is wet. This material MUST be used quickly because there is a gypsum (plaster) binder present.

4. Spread the mix by using a medicine dropper. Do not use disposable pipets! The disposable pipets have extremely narrow openings at the end and they clog up easily. There exists a “dipping method” for preparing TLC slides, but since the usual solvents, methanol and chloroform (Caution! Toxic!) do not activate the binder, the powder falls off the plate. Because the layers formed by this process are very thin, they are very fragile.

5. Run a bead of mix around the outside of the slide, then fill the remaining clear space. Leave ¼ in. of the slide blank on one end, so you can hold onto the slide. Immediately tap the slide from the bottom to smooth the mix out (Fig. 89). Repeat this with as many slides as you can. If the mix sets up and becomes unmanageable, add a little water and shake well.

6. Let the slides sit until the gloss of water on the surface has gone. Then place the slides on a hot plate until they dry. (CAUTION! If the hot plate is too hot, the water will quickly turn to steam and blow the adsorbant off the slides.)

---

**Fig. 89** Spreading adsorbant on a TLC plate.
THE PLATE SPOTTER

1. The spotter is the apparatus used to put the solutions you want to analyze on the plate. You use it to make a spot of sample on the plate.
2. Put the center of a melting point capillary into a small, blue Bunsen burner flame. Hold it there until the tube softens and starts to sag. Do not rotate the tube, ever.
3. Quickly remove capillary from the flame, and pull both ends (Fig. 90). If you leave the capillary in the flame too long, you get an obscene-looking mess.
4. Break the capillary at the places shown in Fig. 90 to get two spotters that look roughly alike. (If you’ve used capillaries with both ends open already, then you don’t have a closed end to break off.)
5. Make up 20 of these or more. You’ll need them.
6. Because TLC is so sensitive, spotters tend to “remember” old samples if you reuse them. Don’t put different samples in the same spotter.

SPOTTING THE PLATES

1. Dissolve a small portion (1–3 mg) of the substance you want to chromatograph in any solvent that dissolves it and evaporates rapidly. Dichloromethane or diethyl ether often works best.

![Fig. 90](image)

**Fig. 90** Making capillary spotters from melting point tubes.
2. Put the thin end of the capillary spotter into the solution. The solution should rise up into the capillary.

3. Touch the capillary to the plate briefly! The compound will run out and form a small spot. Try to keep the spot as small as possible; NOT larger than \( \frac{1}{4} \) in. in diameter. Blow gently on the spot to evaporate the solvent. Touch the capillary to the same place. Let this new spot grow to be almost the same size as the one already there. Remove the capillary and gently blow away the solvent. This will build up a concentration of the compound.

4. Take a sharp object (an old pen point, capillary tube, spatula edge, etc.) and draw a straight line through the adsorbant, as close to the clear glass end as possible (Fig. 91). Make sure that it runs all the way across the end of the slide and goes right down to the glass. This will keep the solvent from running up to the ragged edge of the adsorbant. It will travel only as far as the smooth line you have drawn. Measurements will be taken from this line.

5. Now make a small notch in the plate at the level of the spots to mark their starting position. You'll need this later for measurements.

DEVELOPING A PLATE

1. Take a 150-ml beaker, line the sides of it with filter paper, and cover it with a watch glass (Fig. 92).

2. Choose a solvent to develop the plate. You let this solvent (eluent) pass through the adsorbant by capillary action. Nonpolar eluents (solvents) will force nonpolar compounds to the top of the plate, whereas polar eluents will force BOTH polar and nonpolar materials up the plate. There is only one way to choose eluents. Educated guesswork. Use the chart of eluents in Chapter 18.

3. Pour some of the eluent (solvent) into the beaker, and tilt the beaker so that the solvent wets the filter paper. Put no more than \( \frac{1}{4} \) in. of eluent in the bottom of the beaker!

4. Place the slide into the developing chamber as shown (Fig. 93). Don't let the solvent in the beaker touch the spot on the plate or the spot will dissolve away into the solvent! If this happens, you'll need a new plate, and you'll have to clean the developing chamber as well.
5. Cover the beaker with a watch glass. The solvent (eluent) will travel up the plate. The filter paper keeps the air in the beaker saturated with solvent so that it doesn’t evaporate from the plate. When the solvent reaches the line, immediately remove the plate. Drain the solvent from it, and blow gently on the plate until all the solvent is gone. If not, there will be some trouble visualizing the spots.

Don’t breathe fumes of the eluents! Make sure you have adequate ventilation. Work in a hood if possible.

Fig. 91 Putting a spot of compound on a TLC plate.

Fig. 92 The secret identity of a 150-ml beaker as a TLC slide development chamber is exposed.
VISUALIZATION

Unless the compound is colored, the plate will be blank, and you won’t be able to see anything, so you must visualize the plate.

1. **Destructive visualization.** Spray the plate with sulfuric acid, then bake in an oven at 110°C for 15–20 min. Any spots of compound will be charred blots, utterly destroyed. All spots of compound will be shown.

2. **Semidestructive visualization.** Set up a developing tank (150-ml beaker) but leave out the filter paper and any solvent. Just a beaker with a cover. Add a few crystals of iodine. Iodine vapors will be absorbed onto most spots of compound, coloring them. Removing the plate from the chamber causes the iodine to evaporate from the plate, and the spots will slowly disappear. Not all spots may be visible. So if there’s nothing there, that doesn’t mean nothing’s there. The iodine might have reacted with some spots, changing their composition. Hence the name semidestructive visualization.

3. **Nondestructive visualization.**
   
a. **Long-wave UV (Hazard!)** Most TLC adsorbants contain a fluorescent powder that glows bright green when under long-wave UV light. There are two ways to see the spots:
(1) The background glows green, the spots are dark.

(2) The background glows green, the spots glow some other color. The presence of excess eluent may cause whole sections of the plate to remain dark. Let all the eluent evaporate from the plate.

b. Short-wave UV (Hazard!) The plates stay dark. Only the compounds may glow. This is usually at 180 nm.

Both the UV tests can be done in a UV light box, in a matter of seconds. Since most compounds are unchanged by exposure to UV, the test is considered nondestructive. Not everything will show up, but the procedure is good enough for most compounds. When using the light box, always turn it off when you leave it. If you don’t, not only does the UV filter burn out, but your instructor becomes displeased.

Since neither the UV nor the iodine test is permanent, it helps to have a record of what you’ve seen. You must draw an accurate picture of the plate in your notebook. Using a sharp-pointed object (pen point, capillary tube, etc.) you can trace the outline of the spots on the plate while they are under the UV light (Caution! Wear gloves!) or before the iodine fades from the plate.

INTERPRETATION

After visualization, there will be a spot or spots on the plate. Here is what you do when you look at them.

1. Measure the distance from that solvent line drawn across the plate to where the spot started.

2. Measure the distance from where the spot stopped to where the spot began. Measure to the center of the spot, rather than to one edge. If you have more than one spot, get a distance for each. If the spots are shaped funny, do your best.

3. Divide the distance the solvent moved into the distance the spot(s) moved. The resulting ratio is called the $R_f$ value. Mathematically, the ratio for any spot should be between 0.0 and 1.0, or you goofed. Practically, spots with $R_f$ values greater than about 0.8 and less than about 0.2 are hard to interpret. They could be single spots or multiple spots all bunched up and hiding behind one another.
4. Check out the $R_f$ value—it may be helpful. In identical circumstances, this value would always be the same for a single compound, all the time. If this were true, you could identify unknowns by running a plate and looking up the $R_f$ value. Unfortunately, the technique is not that good, but you can use it with some judgment and a reference compound to identify unknowns (see “Multiple Spotting,” following).

Figures 94 to 96 provide some illustrations. Look at Fig. 94: if you had a mixture of compounds, you could never tell. This $R_f$ value gives no information. Run this compound again. Run a new plate. Never redevelop an old plate!

**Use a more polar solvent!**

No information in Fig. 95, either. You couldn’t see a mixture if it were here. Run a new plate. *Never redevelop an old plate!*

**Use a less polar solvent!**

If the spot moves somewhere between the two limits (shown in Fig. 96) and remains a single spot, the compound is pure. If more than one spot shows, the compound is impure and it is a mixture. Whether the compound should be purified is a matter of judgment.

![Fig. 94](image) Development with a nonpolar solvent and no usable results.
**Fig. 95** Development with a very polar solvent and no usable results.

**Fig. 96** Development with just the right solvent is a success.
MULTIPLE SPOTTING

You can run more than one spot, either to save time or to make comparisons. You can even identify unknowns.

Let’s say that there are two unknowns, A and B. Say one of them can be biphenyl (a colorless compound that smells like moth balls). You spot two plates. One with A and biphenyl, side by side. The other, B and biphenyl, side by side. After you develop both plates, you have the results shown in Fig. 97.

Apparently A is biphenyl.

Note that the $R_f$ values are not perfect. This is an imperfect world, so don’t panic over a slight difference.

And now we have a method that can quickly determine:

![Figure 97](image)

**Fig. 97** Side-by-side comparison of an unknown and a leading brand known.
1. Whether a compound is a mixture.
2. The identity of a compound if a standard is available.

There are preprepared plates having an active coating on a thin plastic sheet, also with or without the fluorescent indicator. You can cut these to any size (they are about 8 in. by 8 in.) with a pair of scissors. Don’t touch the active surface with your fingers — handle them only by the edge. The layers on the plate are much thinner than those you would make by spreading adsorbent on a microscope slide, so you have to use smaller amounts of your compounds so you don’t overload the adsorbent.

As many people have taken the time to point out, you can substitute a much less expensive wide-mouth screw cap bottle for a beaker as a developing chamber.

PREPARATIVE TLC

When you use an analytical technique (like TLC) and you expect to isolate compounds, it’s often called a preparative (prep) technique. So TLC becomes “prep TLC.” You use the same methods only on a larger scale.

Instead of a microscope slide, you usually use a 12 × 12-in. glass plate and coat it with a thick layer of adsorbant (0.5 – 2.0 mm). Years ago, I used a small paintbrush to put a line (a streak rather than a spot) across the plate near the bottom. Now you can get special plate streakers that give a finer line and less spreading. You put the plate in a large developing chamber and develop and visualize the plate as usual.

The thin line separates and spreads into bands of compounds, much like a tiny spot separates and spreads on the analytical TLC plates. Rather than just look at the bands though, you scrape the adsorbant holding the different bands into different flasks, blast your compounds off of the adsorbants with appropriate solvents, filter off the adsorbant, and finally evaporate the solvents and actually recover the separate compounds.
Wet-Column Chromatography
This is, as you may have guessed, chromatography carried out on a column of adsorbant, rather than a layer. Not only is it cheap, easy, and carried out at room temperature but you can separate large amounts, gram quantities, of mixtures.

In column chromatography, the adsorbant is usually alumina but can be silica gel. Except that alumina tends to be basic and silica gel, acidic, I don’t know why the former is used more often. Remember, if you try out an eluent (solvent) on silica gel plates, the results on an alumina column may be different.

Now you have a glass tube as the support holding the adsorbant alumina in place. You dissolve your mixture and put it on the adsorbant at the top of the column. Then you wash the mixture down the column using at least one eluent (solvent), perhaps more. The compounds carried along by the solvent are washed entirely out of the column, into separate flasks. Then you isolate the separate fractions.

**PREPARING THE COLUMN**

1. The alumina is supported by a glass tube with either a stopcock or a piece of tubing and a screw clamp to control the flow of eluent (Fig. 98). You can use an ordinary buret. What you will use will depend on your own lab program. Right above this control you put a wad of cotton or glass wool to keep everything from falling out. Do not use too much cotton or glass wool, and do not pack it too tightly. If you ram the wool into the tube, the flow of eluent will be very slow, and you’ll be in lab till next Christmas waiting for the eluent. If you pack it too loosely, all the stuff in the column will fall out.

2. At this point, fill the column half-full with the least polar eluent you will use. If this is not given, you can surmise it from a quick check of separation of the mixture on a TLC plate. This would be the advantage of an alumina TLC plate.

3. Slowly put sand into the column through a funnel until there is a 1-cm layer of sand over the cotton. Adsorbant alumina is SO FINE, it is likely to go through cotton or glass wool but NOT through a layer of fine sand.
4. During this entire procedure, keep the level of the solvent above that of any solid material in the column!

5. Now slowly add the alumina. Alumina is an adsorbant and it sucks up the solvent. When it does, heat is liberated. The solvent may boil and ruin the column. Add the alumina slowly! Use about 25 g of alumina for every 1 g
of mixture you want to separate. While adding the alumina, tap or gently swirl the column to dislodge any alumina or sand on the sides. You know, a plastic wash bottle with eluent in it can wash the stuff down the sides of the column very easily.

6. When the alumina settles, you normally have to add sand (about 1 cm) to the top to keep the alumina from moving around.

7. Open the stopcock or clamp and let solvent out until the level of the solvent is just above the upper level of sand.

8. Check the column! If there are air bubbles or cracks in the column of alumina, dismantle the whole business and start over!

**COMPOUNDS ON THE COLUMN**

If you’ve gotten this far, congratulations! Now you have to get your mixture, the analyte, on the column. Dissolve your mixture in the same solvent you are going to put through the column. Try to keep the volume of the solution of mixture as small as possible. If your mixture does not dissolve entirely, and it is important that it do so, check with your instructor! You might be able to use different solvents for the analyte and for the column, but this isn’t as good. You might use the least polar solvent that will dissolve your compound.

If you must use the column eluent as the solvent, and not all the solvent will dissolve, you can filter the mixture through filter paper. Try to keep the volume of solution down to 10 ml or so. After this, the sample becomes unmanageable.

1. Use a pipet and rubber bulb to slowly and carefully add it to the top of the column (Fig. 99). Do not disturb the sand!

2. Open the stopcock or clamp and let solvent flow out until the level of the solution of compound is slightly above the sand. At no time let the solvent level get below the top layer of sand! The compound is now “on the column.”

3. Now add eluent (solvent) to the column above the sand. Do not disturb the sand! Open the stopcock or clamp. Slowly let eluent run through the column until the first compound comes out. Collect the different products in Erlenmeyer flasks. You may need lots and lots of Erlenmeyer
flasks. *At no time let the level of the solvent get below the top of the sand!* If necessary, stop the flow, add more eluent, and start the flow again.

**VISUALIZATION AND COLLECTION**

If the compounds are colored, you can watch them travel down the column and separate. If one or all are colorless, you have problems. So:

1. *Occasionally* let 1 or 2 drops of eluent fall on a clean glass microscope slide. Evaporate the solvent and see if there is any sign of your crystalline compound! This is an excellent spot test, but don’t be confused by nasty plasticizers from the tubing trying to put one over on you, pretending to be your product.

2. Put the narrow end of a “TLC spotter” to a drop coming off of the column. The drop will rise up into the tube. Using this loaded spotter, *spot, develop, and visualize a TLC plate with it.* Not only is this more
sensitive, but you can see whether the stuff coming out of the column is pure (see Chapter 19, "Thin-Layer Chromatography"). You'll probably have to collect more than one drop on a TLC plate. If it is very dilute, the plate will show nothing, even if there actually is compound there. It is best to sample 4 or 5 consecutive drops.

Once the first compound or compounds have come out of the column, those that are left may move down the column much too slowly for practical purposes. Normally you start with a nonpolar solvent. But by the time all the compounds have come off, it may be time to pick up your degree. The solvent may be too nonpolar to kick out later fractions. So you have to decide to change to a more polar solvent. This will kick the compound right out of the column.

To change solvent in the middle of a run:

1. Let old solvent level run down to just above the top of the sand.
2. Slowly add new, more polar solvent and do not disturb the sand.

You, and you alone, have to decide if and when to change to a more polar solvent. (Happily, sometimes you'll be told.)

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**Fig. 100** A growth of crystals occurs as the eluent evaporates.
If you have only two components, start with a nonpolar solvent, and when you are sure the first component is completely off the column, change to a really polar one. With only two components, it doesn’t matter what polarity solvent you use to get the second compound off the column.

Sometimes the solvent evaporates quickly and leaves behind a “fuzz” of crystals around the tip (Fig. 100). Just use some fresh solvent to wash them down into the collection flask.

Now all the components are off the column and in different flasks. Evaporate the solvents (No flames!), and lo! The crystalline material is left.

Dismantle the column. Clean up. Go home.
Dry-column chromatography is another approach to the separation of large quantities of a mixture of products. I think it’s easier than wet-column chromatography, though more limited.

1. Weigh about 80 g of the adsorbant (alumina, silica gel, etc.) for a typical \(15 \times \frac{3}{4}\)-in. column, into either a large beaker or a large screw-cap bottle.

2. Get some of your eluent and prewet the adsorbant somewhat. For about 10 g of adsorbant, start with 3–5 g of the liquid eluent. (Oh. Did anyone ever tell you, you can weigh liquids directly, just like solids?) Now add the eluent to the adsorbant and mix like mad. You can see the advantage of a screw-cap bottle over an open beaker. The powder can’t fly out of a closed bottle. Do not add too much eluent! You only want to precondition the adsorbant so that you don’t get bubbling in the column from the heat of hydration released when you eventually run the experiment.

   **The powder should still flow as a powder.**

3. Get a length of flexible, flat nylon tubing. Fold one end over several times (Fig. 101) and staple it to close it off.

4. Open the other end and add sand until the bottom is full (about 2–5 cm).

5. Add the prewet, “dry” adsorbant.

6. Cover with a layer of sand (about 1 cm).

7. Gently clamp this nylon sausage into an upright position (Fig. 101).

8. The manufacturers of dry-column chromatography adsorbants suggest piercing the bottom of the column with a few pin holes. They would know.

9. Use a disposable pipet to load your sample onto the column. The sample should go through the sand and become stuck on the adsorbant.

10. Again, with a disposable pipet, carefully add clean eluent until you’re sure all the sample is stuck on the adsorbant, and none of the sand.

11. At this point, begin carefully adding more eluent. As the eluent goes down the column, the compounds in your mixture will also travel down the column at different rates, and you should get a separation. It is not good to keep adding eluent until it begins to drip out the closed end onto your shoes.

12. When you’re finished, you get to lie the column down carefully on the
benchtop and slice out the sections of the column that have the fractions you want. It's somewhat messy, but just pinch the top of the column closed at the level of the sand, and drain any eluent at the top into a waste container. Probably some sand will go with it. With both ends closed nothing will move around as you move the column.

13. To recover your products, take just that section of the adsorbant with the sample you want on it, put it into a beaker, and wash your product off the adsorbant with a more polar solvent. Then filter off the adsorbant, strip off the solvent, and voilà! Clean, separated material.
NOTE: Be careful with this technique for colorless samples. The reasons are pretty simple, if not obvious. How’re you going to see where to make the cuts if you run colorless samples? Even though nylon tubing is transparent to UV light, and theoretically you can see the compounds under UV, many eluents absorb in the UV, and the whole column would just look dark. Then you’d probably get into trouble guessing where to make the cuts in the tubing to get the compounds separated. Just stick to colored compounds. And don’t say I didn’t warn you.
When light travels from one medium to another it changes velocity and direction a bit. If you’ve ever looked at a spoon in a glass of water, the image of the spoon in water is displaced a bit from the image of the spoon in air, and the spoon looks broken. When the light rays travel from the spoon in water and break out into the air, they are refracted, or shifted (Fig. 102). If we take the ratio of the sine of the angles formed when a light ray travels from air to water, we get a single number, the index of refraction, or refractive index. Because we can measure the index of refraction to a few parts in 10,000, this is a very accurate physical constant for identification of a compound.

The refractive index is usually reported as $n^\circ_{D}$, where the tiny 25 is the temperature at which the measurement was taken, and the tiny capital D means we’ve used light from a sodium lamp, specifically a single yellow frequency called the sodium D line. Fortunately, you don’t have to use a sodium lamp if you have an Abbé refractometer.
The Abbe Refractometer (Fig. 103)

The refractometer looks a bit like a microscope. It has

1. **An eyepiece.** You look in here to make your adjustments and read the refractive index.

2. **Compensation prism adjustment.** Since the Abbe refractometer uses white light and not light of one wavelength (the sodium D line), the white light *disperses* as it goes through the optics and rainbow-like color fringing shows up when you examine your sample. By turning this control, you rotate some compensation prisms that eliminate this effect.

3. **Hinged sample prisms.** This is where you put your sample.

4. **Light source.** This provides light for your sample. It’s on a moveable arm, so you can swing it out of the way when you place your samples on the prisms.

5. **Light source swivel arm lock.** This is a large slotted nut that works itself loose as you move the light source up and down a few times. Always have a dime handy to help you tighten this locking nut when it gets loose.

6. **Sample and scale image adjust.** You use this knob to adjust the optics such that you see a split field in the eyepiece. The refractive index scale also moves when you turn this knob. The knob is often a dual control; use the outer knob for a coarse adjustment and the inner knob as a fine adjustment.

7. **Scale/sample field switch.** Press this switch, and the numbered refractive index scale appears in the eyepiece. Release this switch, and you see your sample in the eyepiece. Some models don’t have this type of switch. You have to change your angle of view (shift your head a bit) to see the field with the refractive index reading.

8. **Line cord on — off switch.** This turns the refractometer light source on and off.

9. **Water inlet and outlet.** These are often connected to temperature-controlled water recirculating baths. The prisms and your samples in the prisms can all be kept at the temperature of the water.
USING THE ABBÉ REFRACTOMETER

1. Make sure the unit is plugged in. Then turn the on-off switch to ON. The light at the end of the moveable arm should come on.

2. Open the hinged sample prisms. NOT touching the prisms at all, place a few drops of your liquid on the lower prism. Then, swing the upper prism back over the lower one and gently close the prisms. Never touch the prisms with any hard object or you’ll scratch them.

3. Raise the light on the end of the moveable arm so that the light illuminates the upper prism. Get out your dime and, with permission of your instructor, tighten the light source swivel arm lock nut as it gets tired and lets the light drop.
4. Look in the eyepiece. Slowly, carefully, with very little force, turn the large scale and sample image adjust knob from one end of its rotation to the other. Do not FORCE! (If your sample is supposedly the same as that of the last person to use the refractometer, you shouldn’t have to adjust this much if at all.)

5. You are looking for a split optical field of light and dark (Fig. 104). This may not be very distinct. You may have to raise or lower the light source and scan the sample a few times.

6. If you see color fringing at the boundary between light and dark (usually red or blue), slowly turn the compensating prism adjust until the colors are at a minimum. You may now have to go back and readjust the sample image knob a bit after you do this.

7. Press and hold the scale/sample field switch. The refractive index scale should appear (Fig. 105). Read the uppermost scale, the refractive index, to four decimal places. (If your model has two fields, with the refractive index always visible, just read it.)

**Fig. 104** Your sample through the lens of the refractor.

**Fig. 105** A refractive index of 1.4398.
REFRACTOMETRY HINTS

1. The refractive index changes with temperature. If your reading is not the same as that of a handbook, check the temperatures before you despair.

2. Volatile samples require quick action. Cyclohexene, for example, has been known to evaporate from the prisms of unthermostatted refractometers more quickly than you can obtain the index. It may take several tries as you readjust the light, turn the sample and scale image adjust, and so on.

3. Make sure the instrument is level. Often organic liquids can seep out of the jaws before you are ready to make your measurement.
Instrumentation in the Lab
Electronic instrumentation is becoming more and more common in the organic lab, which is both good and bad. The good part is that you’ll be able to analyze your products, or unknowns, much faster, and potentially with more accuracy than ever. The bad part is that you have to learn about how to use the instrumentation, and there are many different manufacturers of different models of the same instrument.

The usual textbook approach is to take a piece of equipment, say something like “This is a typical model,” and go on from there, trying to illustrate some very common principles. Only what if your equipment is different? Well, that’s where you’ll have to rely on your instructor to get you out of the woods. I’m going to pick out specific instruments as well. But at least now you won’t panic if the knobs and settings on yours are not quite the same.

With that said, I’d like to point out a few things about the discussions that follow.

1. If you just submit samples to be run on various instruments, as I did as an undergraduate, pay most attention to the sample preparation sections. Often they say not much more than “Don’t hand in a dirty sample,” but often that’s enough.

2. If you get to put the sample into the instruments yourself, sample introduction is just for you.

3. If you get to play with (in a nonpejorative sense) the instrument settings, you’ll have to wade through the entire description.

You’ll notice I’ve refrained from calling these precision instruments “machines.” That’s because they are precision instruments, not machines — unless they don’t work.
Gas chromatography (GC) can also be referred to as vapor-phase chromatography (VPC) and even gas—liquid chromatography (GLC). Usually the technique, the instrument, and the chart recording of the data are all called GC:

"Fire up the GC." (the instrument)
"Analyze your sample by GC." (perform the technique)
"Get the data off the GC." (analyze the chromatogram)

I've mentioned the similarity of all chromatography, and just because electronic instrumentation is used, there's no need to feel that something basically different is going on.

THE MOBILE PHASE: GAS

In column chromatography the mobile (moving) phase is a liquid that carries your material through an adsorbant. I called this phase the eluent, remember? Here a gas is used to push, or carry, your vaporized sample, and it's called the mobile phase.

The carrier gas is usually helium, though you can use nitrogen. You use a microliter syringe to inject your sample into this gas stream through an injection port, then onto the column. If your sample is a mixture, the compounds separate on the column and reach the detector at different times. As each component hits the detector, the detector generates an electric signal. Usually the signal goes through an attenuator network, then out to a chart recorder to record the signal. I know, it's a fairly general description, and Fig. 106 is a highly simplified diagram, but there are lots of different GCs, so being specific about their operation doesn't help here. You should see your instructor. But that doesn't mean we can't talk about some things.

GC SAMPLE PREPARATION

Sample preparation for GC doesn't require much more work than handing in a sample to be graded. Clean and dry, right? Try to take care that the boiling
The sample enters the GC at the injection port (Fig. 107). You use a micro-liter syringe to pierce the rubber septum and inject the sample onto the column. Don’t stab yourself or anyone else with the needle. Remember, this is not dart night at the pub. Don’t throw the syringe at the septum. There is a way to do this.

1. To load the sample, put the needle into your liquid sample and slowly pull the plunger to draw it up. If you move too fast, and more air than sample gets in, you’ll have to push the plunger back again and draw it up once more. Usually they give you a 10-μl syringe, and 1 maybe 2μl of sample is enough. Take the loaded syringe out of the sample, and carefully, cautiously pull the plunger back so there is no sample in the needle. You should see a bit of air at the very top, but not very much. This way, you
Fig. 107 A GC injection port.

don’t run the risk of having your compound boil out of the needle as it enters the injector oven just before you actually inject your sample. That makes the sample broaden and reduces the resolution. In addition, the air acts as an internal standard. Since air travels through the column almost as fast as the carrier gas, the air peak that you get can signal the start of the chromatogram, much like the notch at the start of a TLC plate. Ask your instructor.

2. Hold the syringe in two hands (Fig. 108). There is no reason to practice being an M.D. in the organic laboratory.

3. Bring the syringe to the level of the injection port, straight on. No angles. Then let the needle touch the septum at the center.

4. The real tricky part is holding the barrel and, without injecting, pushing the needle through the septum. This is easier to write about than it is to do the first time.

5. Now quickly and smoothly push on the plunger to inject the sample, and pull the syringe needle out of the septum and injection port.

After a while, the septum gets full of holes and begins to leak. Usually, you can tell you have a leaky septum when the pen on the chart recorder wanders about aimlessly without any sample injected.
Fig. 108 Three little steps to a great GC.

SAMPLE IN THE COLUMN

Now that the sample is in the column, you might want to know what happens to this mixture. Did I say mixture? Sure. Just as with thin-layer and column
chromatography, you can use GC to determine the composition or purity of your sample.

Let's start with two components, A and B again, and follow their path through an adsorption column. Well, if A and B are different, they are going to stick on the adsorbant to different degrees and spend more or less time flying in the carrier gas. Eventually, one will get ahead of the other. Aha! Separation — Just like column and thin-layer chromatography. Only here the samples are vaporized, and it's called vapor-phase chromatography (VPC).

Some of the adsorbants are coated with a liquid phase. Most are very high-boiling liquids, and some look like waxes or solids at room temperature. Still, they're liquid phases. So, the different components of the mixture you've injected will spend different amounts of time in the liquid phase and, again, a separation of components in your compound. Thus the technique is known as gas–liquid chromatography (GLC). Thus you could use the same adsorbant and different liquid phases, and change the characteristics of each column. Can you see how the sample components would partition themselves between the gas and liquid phases and separate according to, perhaps, molecular weight, polarity, size, and so on, making this technique also known as liquid-partition chromatography?

Since these liquid phases on the adsorbant are, eventually, liquids, you can boil them. And that's why there are temperature limits for columns. It is not the best to heat a column past the recommended temperature, boiling the liquid phase right off the adsorbant and right out of the instrument.

High temperature and air (oxygen) are death for some liquid phases, since they oxidize. So make sure the carrier gas is running through them at all times, even a tiny amount, while the column is hot.

**SAMPLE AT THE DETECTOR**

There are several types of detectors, devices that can tell when a sample is passing by them. They detect the presence of a sample and convert it to an electrical signal that's turned into a GC peak (Fig. 109) on the chart recorder. The most common type is the thermal conductivity detector. Sometimes called "hot-wire detectors," these devices are very similar to the filaments you
### Table: GC Trace Measurements

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value 1</th>
<th>Value 2</th>
<th>Value 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Height (mm)</td>
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<td>68</td>
<td>145</td>
</tr>
<tr>
<td>Width at half-height (mm)</td>
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<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Area (mm²)</td>
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<td>580</td>
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<tr>
<td>Relative area</td>
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<td>8.5</td>
<td>18.1</td>
</tr>
<tr>
<td>Distance from injection (in.)</td>
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<td>3¾₁₆</td>
<td>3¹₁₄₁₆</td>
</tr>
<tr>
<td>Retention times $t_R$ at 2 min/in. (min)</td>
<td>6</td>
<td>6.63</td>
<td>7.38</td>
</tr>
</tbody>
</table>

![GC Trace Diagram](image)

**Fig. 109** A well-behaved GC trace showing a mixture of three compounds.
find in light bulbs, and they require some care. Don’t ever turn on the filament current unless the carrier gas is flowing. A little air (oxygen), a little heat, a little current, and you get a lot of trouble replacing the burned-out detector.

Usually there’ll be at least two thermal conductivity detectors in the instrument, in a “bridge circuit.” Both detectors are set in the gas stream, but only one gets to see the samples. The electric current running through them heats them up, and they lose heat to the carrier gas at the same rate.

As long as no samples, only carrier gas, goes over both detectors, the bridge circuit is balanced. There’s no signal to the recorder, and the pen does not move.

Now a sample in the carrier gas goes by one detector. This sample has a thermal conductivity different from that of pure carrier gas. So the sample detector loses heat at a different rate from the reference detector. (Remember, the reference is the detector that NEVER sees samples — only carrier gas.) The detectors are in different surroundings. They are not really equal any more. So the bridge circuit becomes unbalanced and a signal goes to the chart recorder, giving a GC peak.

Try to remember the pairing of sample with reference and that it’s the difference in the two that most electronic instrumentation responds to. You will see this again and again.

**ELECTRONIC INTERLUDE**

There are two other stops the electrical signal makes on its way to the chart recorder.

1. **The coarse attenuator.** This control makes the signal weaker (attenuates it). Usually there’s a scale marked in powers of two: 2, 4, 8, 16, 32, 64, … So each position is half as sensitive as the last one. There is one setting, either an ∞ or an S (for shorted), which means that the attenuator has shorted out the terminals connected to the chart recorder. Now the chart recorder zero can be set properly.

2. **The GC Zero control.** This is a control that helps set the zero position on the chart recorder, but it is not to be confused with the zero control on the chart recorder.
Here's how to set up the electronics, properly, for a GC and a chart recorder.

1. The chart recorder and GC should be allowed to warm up and stabilize for at least 10–15 min. Some systems take more time; ask your instructor.
2. Set the coarse attenuator to the highest attenuation, usually an $\infty$ or S (for shorted).
3. Now set the pen on the chart recorder to zero using only the chart recorder zero control. Once you do that, leave the chart recorder alone.
4. Start turning the coarse attenuator control to more sensitive settings (lower numbers) and watch the pen on the chart recorder.
5. If the pen on the chart recorder moves off zero, use the GC zero control only to bring the pen back to the zero line on the chart recorder paper.
6. Do not touch the chart recorder zero. Use the GC zero control only.
7. As the coarse attenuator gets to more sensitive settings (lower numbers), it becomes more difficult to adjust the chart recorder pen to zero using only the GC zero control. Do the best you can at the lowest attenuation (highest sensitivity) you can hold a zero steady at.
8. Now, you don't normally run samples on the GC at attenuations of 1 or 2. These settings are very sensitive, and there may be lots of electrical noise—the pen jumps about. The point is, if the GC zero is OK at an attenuation of 1, then when you run at attenuations of 8, 16, 32, and so on, the baseline will not jump if you change attenuation in the middle of the run.

Now that the attenuator is set to give peaks of the proper height, you're ready to go. Just be aware that there may be a polarity switch that can make your peaks shift direction.

**SAMPLE ON THE CHART RECORDER**

Interpreting a GC is about the same as interpreting a TLC plate, so I'll use TLC terms as comparison to show the similarities. Remember the $R_f$ value from TLC? The ratio of the distance the eluent moved to how far the spots of compound moved? Well, distances can be related to times, so the equivalent of $R_f$ in GC is retention time. It's the time it takes the sample to move
through the column minus the time it takes for the carrier gas to move through the column. Remember the part about putting air into the syringe to get an air peak? Well, you can assume that air travels with the carrier gas and doesn’t interact with the column material. So the air peak that shows up on the chart paper can be considered to be the reference point, the “notch,” as it were, that marks the start, just as on the TLC plates.

OK, so you don’t want to use an air peak. Then make a mark on the chart paper as soon as you’ve injected the sample, and use that as the start. Not as good, but it’ll work.

No. You do not need a stopwatch for the retention times. Find out the distance the chart paper crawls in, say, a minute. Then get out your little ruler and measure the distances from the starting point (either air peak or pen mark) to the midpoint of each peak on the baseline (Fig. 109). Don’t be wise and do any funny angles. It won’t help. You’ve got the distances and the chart speed, so you’ve got the retention time. It works out. Trust me.

You can also estimate how much of each compound is in your sample by measuring peak areas. The area under each GC peak is proportional to the amount of material that’s come by the detector in that fraction. You might have to make a few assumptions (e.g., the peaks are truly triangular and each component gives the same response at the detector), but usually it’s pretty straightforward. Multiply the height of the peak by the width at half the height. If this sounds suspiciously like the area of a triangle, you’re on the right track. It’s usually not half the base times the height, however, since sometimes the baseline is not very even, and that measurement is difficult.

PARAMETERS, PARAMETERS

To get the best GC trace from a given column, there are lots of things you can do, simple because there are so many controls that you have. Usually you’ll be told the correct conditions, or they’ll be preset on the GC.

Gas Flow Rate

The faster the carrier gas flows, the faster the compounds are pushed through the column. Because they spend less time in the stationary phase, they don’t
separate as well, and the GC peaks come out very sharp but not well separated. If you slow the carrier gas down too much, the compounds spend so much time in the stationary phase that the peaks broaden and overlap gets very bad. The optimum is, as always, the best separation you need, in the shortest amount of time. Sometimes the manufacturer of the GC recommends ranges of gas flow. Sometimes you're on your own. Most of the time, someone else has already worked it out for you.

**Temperature**

Whether you realize it or not, the GC column has its own heater — the column oven. If you turn the temperature up, the compounds hotfoot it through the column very quickly. Because they spend less time in the stationary phase, they don’t separate as well, and the GC peaks come out very sharp but not well separated. If you turn the temperature down some, the compounds spend so much time in the stationary phase that the peaks broaden and overlap gets very bad. The optimum is, as always, the best separation you need in the shortest amount of time. There are two absolute limits, though.

1. *Too high and you destroy the column.* The adsorbant may decompose, or the liquid phase may boil out onto the detector. Never exceed the recommended maximum temperature for the column material. Don’t even come within 20°C of it just to be safe.

2. *Way too low a temperature, and the material condenses on the column.* You have to be above the dew point of the least volatile material. Not the boiling point. Water doesn’t always condense on the grass — become dew — every day that’s just below 100°C (that’s 212°F, the boiling point of water). Fortunately, you don’t have to know the dew points for your compounds. You do have to know that you don’t have to be above the boiling point of your compounds

Incidentally, the injector may have a separate injector oven, and the detector may have a separate detector oven. Set them both 10 to 20°C
higher than the column temperature. You can even set these above the boiling points of your compounds, since you do not want them to condense in the injection port or the detector, ever. For those with only one temperature control, sorry. The injection port, column, and detector are all in the same place, all in the same oven, and all at the same temperature. The maximum temperature, then, is limited by the decomposition temperature of the column. Fortunately, because of that dew point phenomenon, you really don’t have to work at the boiling points of the compounds either.
HPLC. Is it high-performance liquid chromatography or high-pressure liquid chromatography or something else? It’s probably easier to consider it a delicate blend of wet-column chromatography and gas chromatography (see Chapters 20 and 24, respectively).

Rather than letting gravity pull the solvent through the powdered adsorbant, the liquid is pumped through under pressure. Initially, high pressures [1000–5000 psig (pounds per square inch gauge), i.e., not absolute] were used to push a liquid through a tightly packed solid. But the technique works well at lower pressures (~250 psig), hence the name high-performance liquid chromatography.

From there on, the setup (Fig. 110) resembles gas chromatography very closely. Although a moving liquid phase replaces the helium stream, compounds are put onto a column through an injection port, they separate inside a chromatographic column in the same way as in GC by spending more or less time in a moving liquid now, and the separated compounds pass through a detector. There the amounts of each compound as they go by the detector are turned into electrical signals and displayed on a chart recorder as HPLC peaks that look just like GC peaks. You should also get the feeling that the analysis of these HPLC traces is done in the same manner as GC traces, because it is.

Again, there’s a lot of variety among HPLC systems, so what I say won’t necessarily apply to your system in every respect. But it should help. I’ve based my observations on a Glencoe HPLC unit. It is simple and rugged, performs very well, and uses very common components carried by almost every HPLC supplier. Parts are easy to get.

That’s probably why the company has stopped making this unit. Anything simple and rugged is not likely to need a lot of attention, nor is it likely to go out of fashion, and there’s little profit in that. The individual pieces can be bought from many chromatographic supply houses. Just follow the directions for connections given here.

THE MOBILE PHASE: LIQUID

If you use only one liquid, either neat or as a mixture, the entire chromatogram is said to be isocratic. There are units that can deliver varying solvent compositions over time. These are called gradient elution systems.
Fig. 110 Block diagram of an HPLC setup.
For an isochratic system, you usually use a single solution, or a neat liquid, and put it into the solvent reservoir, generally a glass bottle with a stopcock at the bottom to let the solvent out (Fig. 111). The solvent travels out of the bottom of the reservoir and usually through a solvent filter that traps out any fairly large, insoluble impurities that may be in the solvent.

It is important, if you're making up the eluent yourself, to follow the directions scrupulously. Think about it. If you wet the entire system with the wrong eluent, you can wait a very long time for the correct eluent to reestablish the correct conditions.

A Bubble Trap

Air bubbles are the nasties in HPLC work. They cause the same type of troubles as with wet-column chromatography, and you just don't want them. So there's usually a bubble trap (Fig. 112) before the eluent reaches the pump. This device is quite simple, really. Bubbles in the eluent stream rise

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**Fig. 111** Aspirator bottle used to deliver eluent.
up the center pipe and are trapped there. To get rid of the bubbles, you open the cap at the top. Solvent then rises in the tube and pushes the bubbles out. You have to be extremely careful about bubbles if you’re the one to start the setup or if the solvent tank has run out. Normally, one bubble purge per day is enough.

The Pump

The most common pumping system is the reciprocating pump. Milton Roy makes a pretty good model. The pump has a reciprocating ruby rod that moves back and forth. On the backstroke, the pump loads up on a little bit of solvent, then it squirts it out, under pressure, on the forward stroke. If you want to increase the amount of liquid going through the system, you can dial the length of the stroke, from zero to a preset maximum, using the micrometer at the front of the pump (see Fig. 113). Use a fully clockwise setting, and the stroke length is zero—no solvent flow. A fully counterclockwise setting gives the maximum stroke length—maximum solvent flow. If you have a chance to work with this type of pump, always turn the micrometer fully clockwise to give a zero stroke length before you start the pump. If there is a stroke length set before you turn the pump on, the first smack can damage the reciprocating ruby rod. And it is not cheap.

The Pulse Dampener

Because the rod reciprocates (i.e., goes back and forth) you’d expect huge swings in pressure, pulses of pressure, to occur. That’s why they make pulse
**dampeners.** A coiled tube is hooked to the pump on the side opposite the column (Fig. 113). It is filled with the eluent that’s going through the system. On the forward stroke, solvent is compressed into this tube and *at the same time*, a shot of solvent is pushed onto the column. On the backstroke, while the pump chamber fills up again, the eluent we just pressed into the pulse dampener *squirts out into the column*. Valves in the pump take care of directing the flow. With the eluent in the pulse dampener tubing taking up the slack, the huge variations in pressure, from essentially zero to perhaps 1000 psig, are evened out. They don’t disappear, going about 100 psig either way, but these ***dampened pulses*** are now too small to be picked up on the detector. They don’t show up on the chart recorder either.
HPLC SAMPLE PREPARATION

Samples for HPLC must be liquids or solutions. It would be nice if the solvent you’ve dissolved your solid sample in were the same as the eluent.

It is absolutely crucial that you preclean your sample. Any decomposed or insoluble material will stick to the top of the column and can continually poison further runs. There are a few ways to keep your column clean.

1. **The Swinney adapter** (Fig. 114). This handy unit locks onto a syringe already filled with your sample. Then you push the sample slowly through a Millipore filter to trap insoluble particles. This does not, however, get rid of soluble tars that can ruin the column. (Oh. Don’t confuse the filters with the papers that separate them. It’s embarrassing.)

2. **The precolumn filter** (Fig. 115). Add a tiny column, filled with exactly the same material as the main column, and let this small column get contaminated. Then unscrew it, clean out the gunk and adsorbant, and

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Fig. 114 The Swinney adapter and syringe parts.
refill it with fresh column packing. The disadvantage is that you don’t really know when the garbage is going to poison the entire precolumn filter and then start ruining the analyzing column. The only way to find out whether you have to clean the precolumn is to take it out of the instrument. You really want to clean it out long before the contaminants start to show up at the precolumn exit.

**HPLC SAMPLE INTRODUCTION**

This is the equivalent of the **injection port** for the GC technique. With GC you could inject through a rubber septum directly onto the column. With HPLC it’s very difficult to inject against a liquid stream moving at possibly 1000 psig. That’s why they invented **injection port valves** for HPLC: you put your sample into an **injection loop** on the valve that is not in the liquid stream, then **turn the valve**, and voilà, your sample is in the stream, headed for the column.

The valve (Fig. 116) has two positions.
1. **Normal solvent flow.** In this position, the eluent comes into the valve, goes around, and comes on out into the column without any bother. *You load the sample loop in this position.*

2. **Sample introduction.** Flipped this way, the eluent is pumped through the sample loop and any sample there is carried along and into the column. *You put the sample on the column in this position.*

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**SAMPLE IN THE COLUMN**

Once the sample is in the column, there’s not much difference between what happens here and what happens in paper, thin-layer, vapor-phase (gas), wet-column, or dry-column chromatography. *The components in the mixture will stay on the stationary phase, or move in the mobile phase for different times and end up at different places when you stop the experiment.*

So what’s the advantage? You can separate and detect microgram quanti-
ties of solid samples much as in GC. And you can’t do solids all that well by GC because you have to vaporize your solid sample, probably decomposing it.

A novel development for HPLC is something called **bonded reversed-phase columns**, where the stationary phase is a nonpolar hydrocarbon, chemically bonded to a solid support. You can use these with aqueous eluents, usually alcohol–water mixtures. So you have a **polar eluent and a nonpolar stationary phase**, something that does not usually occur for ordinary wet-column chromatography. One advantage is that you don’t need to use anhydrous eluents (very small amounts of water can change the character of normal phase columns) with reversed-phase columns.

## SAMPLE AT THE DETECTOR

There are many HPLC detectors that can turn the presence of your compound into an electrical signal to be written on a chart recorder. Time was the **refractive index** detector was common. Clean eluent, *used as a reference*, went through one side of the detector, and the *eluent with the samples* went through the other side. A **difference in the refractive index between the sample and reference caused an electrical signal to be generated and sent to a chart recorder**. If you’ve read the section on gas chromatography and looked ahead at infrared, you shouldn’t be surprised to find both a **sample** and a **reference**. I did tell you the reference/sample pair is common in instrumentation.

More recently, **UV detectors** (Fig. 117) have become more popular. UV radiation is beyond the purple end of the rainbow, the energy from which great tans are made. So, if you set up a **small mercury vapor lamp** with a power supply to light it up, you’ll have a **source of UV light**. It’s usually filtered to let through only the wavelengths of 180 and/or 254 nm. (And where have you met *that* number before? TLC plates, maybe?) This UV light then passes through a **flow cell** that has the *eluent and your separated sample flowing through it against air as a reference*. **When your samples come through the cell, they absorb the UV, and an electrical signal is generated**. Yes, the signal goes to a chart recorder and shows up as HPLC peaks.
PARAMETERS, PARAMETERS

To waste jar

Separated sample in

Flow cell

Detector

Dual UV beams
Reference is air

Filter passes UV, blocks visible blue glow

Mercury lamp

All glass is quartz, which passes UV,
KEEP GLASS CLEAN!

Fig. 117 Cutaway view of a HPLC UV detector.

SAMPLE ON THE CHART RECORDER

Go back and read about HPLC peak interpretation in the section on GC peak interpretation ("Sample on the Chart Recorder"). The analysis is *exactly the same*, retention times, peak areas, baselines, . . . all that.

PARAMETERS, PARAMETERS

To get the best LC trace from a given column, there are lots of things you can do, most of them the same as for GC (see "Gas Chromatography, Parameters, Parameters").
Eluent Flow Rate

The faster the eluent flows, the faster the compounds are pushed through the column. Because they spend less time in the stationary phase, they don't separate as well and the LC peaks come out very sharp but not well separated. If you slow the eluent down too much, the compounds spend so much time in the stationary phase that the peaks broaden and overlap gets very bad. The optimum is, as always, the best separation you need, in the shortest amount of time. One big difference in LC is the need to worry about back-pressure. If you try for very high flow rates, the LC column packing tends to collapse under the pressure of the liquid. This, then, is the cause of the back-pressure, resistance of the column packing. If the pressures get too high, you may burst the tubing in the system, damaging the pump, ... , all sorts of fun things.

Temperature

Not many LC setups have ovens for temperatures like those for GC. This is because eluents tend to boil at temperatures much lower than the compounds on the column, which are usually solids anyway. And eluent bubbling problems are bad enough, without actually boiling the solvent in the column. This is not to say that LC results are independent of temperature. They're not. But if a column oven for LC is present, its purpose more likely is to keep stray drafts and sudden chills away than to have a hot time.

Eluent Composition

You can vary the composition of the eluent (mobile phase) in HPLC a lot more than in GC, so there's not really much correspondence. Substitute nitrogen for helium in GC and usually the sensitivity decreases, but the retention times stay the same. Changing the mobile phases — the gases — in GC doesn't have a very big effect on the separation or retention time.

There are much better parallels to HPLC: TLC or column chromatography. Vary the eluents in these techniques and you get widely different results. With a normal-phase silica-based column, you can get results similar to those from silica gel TLC plates.
Unlike the chromatographies, which physically separate materials, infrared (IR) spectroscopy is a method of determining what you have after you’ve separated it.

The IR spectrum is the name given to a band of frequencies between 4000 and 650 cm$^{-1}$ beyond the red end of the visible spectrum. The units are called wave numbers or reciprocal centimeters (that’s what cm$^{-1}$ means). This range is also expressed as wavelengths from 2.5 to 15 micrometers ($\mu$m).

With your sample in the sample beam, the instrument scans the IR spectrum. Specific functional groups absorb specific energies. And because the spectrum is laid out on a piece of paper, these specific energies become specific places on the chart.

Look at Fig. 118. Here’s a fine example of a pair of alcohols if ever there was one. See the peak (some might call it trough) at about 3400 cm$^{-1}$ (2.9 $\mu$m)? That’s due to the OH group, specifically the stretch in the O–H bond, the OH stretch.

Now, consider a couple of ketones, 2-butanone and cyclohexanone (Fig. 119). There’s no OH peak about 3400 cm$^{-1}$ (2.9 $\mu$m), is there? Should there be? Of course not. Is there an OH in 2-butanone? Of course not. But there is a C=O, and where’s that? The peak about 1700 cm$^{-1}$ (5.9 $\mu$m). It’s not there for the alcohols and it is there for the ketones. Right. You’ve just correlated or interpreted four IRs.

Because the first two (Fig. 118) have the characteristic OH stretch of alcohols, they might just be alcohols. And the other two (Fig. 119) might be ketones because of the characteristic C=O stretch at 1700 cm$^{-1}$ (5.9 $\mu$m) in each.

What about all the other peaks? You can ascribe some sort of meaning to each of them, but it can be very difficult. That’s why frequency correlation diagrams, or IR tables, exist (Fig. 120). They identify regions of the IR spectrum where peaks for various functional groups show up. They can get very complicated. Check to see if you can find the C–H stretch and the C–O stretch that are in all four spectra, using the correlation table. It can be fun.

For you Sherlock Holmes fans, the region from 1400 to 990 cm$^{-1}$ (7.2–11.1 $\mu$m) is known as the fingerprint region. The peaks are due to the entire molecule, its fingerprint, rather than being from independent functional groups. And, you guessed it, no two fingerprints are alike.

Take another look at the cyclohexanol and cyclohexanone spectra. Both
Fig. 118  IR spectra of (a) t-butanol and (b) cyclohexanol.
Fig. 119 IR spectra of (a) 2-butanone and (b) cyclohexanone.
Approximate infrared absorption frequencies of various groups

- CH
- CH₂
- CH₃
- C = C
- C = C_O_ester
- C = C_Co_olefinic
- C = C_Co_trans
- C = C_Co_cis
- Substitution patterns
- H = C
- C = O
- C = O_Ester
- C = O_Acid
- C = O_Anhydride
- C = O_Acid chloride
- C = C_Amide
- C = C_Aldehydes and ketones
- C = C_NH amides
- C = N_NH amines
- C = N_NH amides

Fig. 120 The author's only IR correlation chart.
have very different functional groups. Now look at the similarities, the simplicity, including the fingerprint region. Both are six-membered rings and have a high degree of symmetry. You should be able to see the similarities due to the similar structural features.

Two more things. Watch your spelling and pronunciation. It’s not “in-fared,” OK? And most people I know use IR (pronounced “eye-are,” not “ear”) to refer to the technique, the instrument, and the chart recording of the spectrum:

"That’s a nice new IR you have there."  (the instrument)
"Take an IR of your sample."  (perform the technique)
"Let’s look at your IR and see what kind of compound you have."  (interpret the resulting spectrogram)

To take an IR, you need an IR. These are fairly expensive instruments; again, no one is typical, but you can get a feeling of how to run an IR as you go on.

**INFRARED SAMPLE PREPARATION**

You can prepare samples for IR spectroscopy easily, but you must strictly adhere to one rule:

**No water!**

In case you didn’t get that the first time:

**No water!**

Ordinarily, you put the sample between two salt plates. Yes. Common, ordinary water-soluble salt plates. Or mix it with potassium bromide (KBr), another water-soluble salt.

So keep it dry, people.
Liquid Samples

1. Make sure the sample is DRY. NO WATER!
2. Put some of the dry sample (2–3 drops) on one plate, then cover it with another plate (Fig. 121). The sample should spread out to cover the entire plate. You don’t have to press. If it doesn’t cover well, try turning the top plate to spread the sample, or add more sample.
3. Place the sandwich in the IR salt plate holder and cover it with a hold-down plate.
4. Put at least two nuts on the posts of the holder (opposite corners) and spin them down GENTLY to hold the plates with an even pressure. Do not use force! You’ll crack the plates! Remember, these are called salt plate holders and not salt plate smashers.
5. Slide the holder and plate into the bracket on the instrument in the sample beam (closer to you, facing instrument).
6. Run the spectrum.

Since you don’t have any other solvents in there, just your liquid com-
pound, you have just prepared a liquid sample, **neat**, meaning *no solvent.* This is the same as a **neat liquid sample**, which is a way of describing any liquid without a solvent in it. It is not to be confused with “really neat liquid sample,” which is a way of expressing your true feelings about your sample.

**Solid Samples**

**The Nujol Mull**

A rapid, inexpensive way to get an IR of solids is to mix them with Nujol, a commercially available mineral oil. Traditionally this is called “making a Nujol mull,” and it is practically idiomatic among chemists. Though you won’t see Rexall or Johnson & Johnson mulls, the generic brand **mineral oil mull** is often used.

You want to disperse the solid throughout the oil, making the solid transparent enough to IR that the sample will give a usable spectrum. Since mineral oil is a saturated hydrocarbon, it has an IR spectrum all its own. You’ll find hydrocarbon bends, stretches, and push-ups in the spectrum, but you know where they are, and you ignore them. You can either look at a published reference Nujol spectrum (Fig. 122) or run your own if you’re not sure where to look.

1. Put a small amount of your solid into a tiny agate mortar and add a few drops of mineral oil.
2. Grind the oil and sample together until the solid is a fine powder *dispersed throughout the oil*.
3. Spread the mull on one salt plate and cover it with another plate. There should be no air bubbles, just an even film of the solid in the oil.
4. Proceed as if this were a **liquid sample**.
5. Clean the plates with *anhydrous* acetone or ethanol. *NOT WATER!* If you don’t have the tiny agate mortar and pestle, try a Witt spot plate and the rounded end of a thick glass rod. The spot plate is a piece of glazed porcelain with dimples in it. Use one as a tiny mortar; the other as a tiny pestle.

And remember to forget the peaks from the Nujol itself.
Fig. 122 A published reference Nujol spectrum.
Solid KBr Methods

KBr methods (hardly ever called potassium bromide methods) consist of making a mixture of your solid (dry again) with IR-quality KBr. Regular KBr off the shelf is likely to contain enough nitrate, as KNO₃, to give spurious peaks, so don’t use it. After you have opened a container of KBr, dry it and later store it in an oven, with the cap off, at about 110°C to keep the moisture out.

Preparing the Solid Solution

1. At least once in your life, weigh out 100 mg of KBr so you’ll know how much that is. If you can remember what 100 mg of KBr looks like, you won’t have to weigh it out every time you need it for IR.
2. Weigh out 1–2 mg of your dry, solid sample. You’ll have to weigh out each sample because different compounds take up different amounts of space.
3. Pregrind the KBr to a fine powder about the consistency of powdered sugar. Don’t take forever, since moisture from the air will be coming in. Add your compound. Grind together. Serves one.

Pressing a KBr Disk — The Mini-Press (Fig. 123)

1. Get a clean, dry press and two bolts. Screw one of the bolts into the press about halfway and call that the bottom of the press.
2. Scrape a finely ground mixture of your compound (1–2 mg) and KBr (approx. 100 mg) into the press so that an even layer covers the bottom bolt.

3. Take the other bolt and turn it in from the top. Gently tighten and loosen this bolt at least once to spread the powder evenly on the face of the bottom bolt.

4. Hand tighten the press again, then use wrenches to tighten the bolts against each other. Don’t use so much force that you turn the heads right off the bolts.

5. Remove both bolts. A KBr pellet, containing your sample, should be in the press. Transparent is excellent. Translucent will work. If the sample is opaque, you can run the IR, but I don’t have much hope of your finding anything.

6. Put the entire press in a holder placed in the analyzing beam of the IR, as in Fig. 124. (Don’t worry about that yet, I’ll get to it in a moment. “Running the Spectrum,” is next.)

Pressing a KBr Disk—The Hydraulic Press

If you have a hydraulic press and two steel blocks available, there is another easy KBr method. It’s a card trick, and at no time do my fingers leave my hands. The only real trick is you’ll have to bring the card.
1. Cut and trim an index card so that it fits into the sample beam aperture (Fig. 129).
2. Punch a hole in the card with a paper punch. The hole should be centered in the sample beam when the card is in the sample beam aperture.
3. Place one of the two steel blocks on the bottom jaw of the press.
4. Put the card on the block.
5. Scrape your KBr-sample mixture onto the card, covering the hole and some of the card. Spread it out evenly.
6. Cover the card, which now has your sample on it, with the second metal block (Fig. 125).
7. Pump up the press to the indicated safe pressure.
8. Let this sit for a bit (1 min). If the pressure has dropped, bring it back up, slowly, carefully, to the safe pressure line, and wait another bit (1 min).
10. Open the metal sandwich. Inside will be a file card with a KBr window in it, just like in the Mini-Press. (CAUTION! The KBr window you form is rather fragile, so don’t beat on it.
11. Put the card in the sample beam aperture (Fig. 126). The KBr win-

![Fig. 125](image)

KBr disks by hydraulic press.
Sample holder plate slips down into groove

Reference aperture

Sample aperture

**Fig. 126** Putting sample holders with samples into the beam.

dow should be *centered* in the sample beam if you’ve cut and punched the card correctly. Now you’re ready to run the spectrum.

**RUNNING THE SPECTRUM**

There are many IR instruments, and since they are so different, you need your instructor’s help here more than ever. But there are a few things you have to know.

1. *The sample beam*. Most IRs are dual-beam instruments (Fig. 127).
Pen traces percent transmission on paper at various wavelengths.

Motor turns grating; drives chart at the same time.

Rotating chopper mirror switches sample and reference beams to detector for comparison.

**Fig. 127** Schematic diagram of an IR.

The one closest to you, if you’re operating the instrument, is the **sample beam**. Logically, there is a **sample holder** for the sample beam, and your sample goes there. And there, a beam of IR radiation goes through your sample.

2. **The reference beam.** This is the other light path. It’s not *visible* light but another part of the electromagnetic spectrum. Just remember that the reference beam is the one farthest away from you.

3. **The 100% control.** This sets the pen at the 100% line on the chart paper. Or tries to. It’s a very delicate control and doesn’t take kindly to excessive force. Read on and all will be made clear.

4. **The pen.** There is a pen and pen holder assembly on the instrument. This is how the spectrum gets recorded on the chart paper. Many people get the urge to throw the instrument out the window when the pen stops writing in the middle of the spectrum. Or doesn’t even start writing. Or was left empty by the last fellow. Or was left to dry out on the top of the instrument. For those with Perkin–Elmer 137s or 710s, two clever fellows have made up generic felt-tip pen holders for the instruments. This way, you buy your own pen from the bookstore, and if it dries out it’s your

5. **The very fast or manual scan.** To get a good IR, you’ll have to be able to scan, very rapidly, *without letting the pen write on the paper*. This is so you’ll be able to make adjustments before you commit pen to paper. This fast forward is not a standard thing. Sometimes you operate the instrument by hand, pushing or rotating the paper holder. Again, *do not use force*.

Whatever you do, *don’t try to move the paper carrier by hand when the instrument is scanning a spectrum*. Stripped gears is a crude approximation of what happens. So, thumbs off.

I’m going to apply these things in the next section using a real instrument, the Perkin–Elmer 710B, as a model. Just because you have another model is no reason to skip this section. If you do have a different IR, try to find the similarities between it and the Perkin–Elmer model. Ask your instructor to explain any differences.

**THE PERKIN–ELMER 710B IR (Fig. 128)**

1. **On-off switch and indicator.** Press this once, the instrument comes on, and the switch lights up. Press this again, the instrument goes off, and the light goes out.

2. **Speed selector.** Selects speed (normal or fast). “Fast” is faster, but slower gives *higher resolution*, that is, more detail.

3. **Scan control.** Press this to *start a scan*. When the instrument is scanning, the optics and paper carrier move automatically, causing the IR spectrum to be drawn on the *chart paper* by the pen.

4. **Chart paper carriage.** This is where the chart paper nestles while you run an IR. If it looks suspiciously like a clipboard, it’s because that’s how it works.

5. **Chart paper hold-down clip.** Just like a clipboard, this holds the paper down in the carrier.

6. **Frequency scale.** This scale is used to help align the chart paper and to tell you during the run where in the spectrum the instrument is.
7. **Scan position indicator.** A white arrow that points, roughly, to the place in the spectrum the instrument is at.

8. **Line-up mark.** A line, here at the number 4000, that you use to match up the numbers on the instrument frequency scale with the same numbers on the chart paper.

9. **Pen and transmittance scale.** This is where the pen traces your IR spectrum. The numbers here mean percentage of IR transmitted through your sample. If you have no sample in the sample beam, how much of the light is getting through? Those who said 100% are 100% correct. Block the beam with your hand and 0% gets through. You should be able to see why these figures are called the % transmission, or %T, scale.

10. **The 100% control.** Sets the pen at 100%. Or tries to. This is a fairly sensitive control, so don’t force it.
11. **Sample beam aperture.** This is where you put the holder containing your sample, be it mull or KBr pellet. You slip the holder into the aperture window for analysis.

12. **Reference beam aperture.** This is where nothing goes. Or, in extreme cases, you use a reference beam attenuator to cut down the amount of light reaching the detector.

### USING THE PERKIN-ELMER 710B

1. Turn the instrument on and let it warm up for about 3–5 min. Other instruments may take longer.

2. Get a piece of IR paper and load the chart paper carriage, just like a clipboard. Move the paper to get the index line on the paper to line up with the index line on the instrument. It’s at 4000 cm$^{-1}$ and it’s only a rough guide. Later I’ll tell you how to calibrate your chart paper.

3. Make sure the chart paper carriage is at the high end of the spectrum (4000 cm$^{-1}$).

4. Put your sample in the sample beam. Slide the sample holder with your sample into the sample beam aperture (Fig. 126).

5. What to do next varies for particular cases. Not much, but enough to be confusing in setting things up.

6. Look at where the pen is. Carefully use the 100% control to locate the pen at about 90% mark when the chart (and spectrum) is at the high end (4000 cm$^{-1}$).

### The 100% Control: An Important Aside

Usually there’s not much more to adjusting the 100% control than is coming up in items 7 through 9, unless your sample, by its size alone, reduces the amount of IR reaching the detector. This really shows up if you’ve used the Mini-Press, which has a much smaller opening than that of the opening in the reference beam. So you’re at a disadvantage right from the start. The 100% control mopes around at, sometimes, much less than 40%. That’s terrible, and you have to use a reference beam attenuator (Fig. 129) to equalize the
Gently slide reference beam attenuator and watch pen — stop when pen points to 80-90% T

7. OK, at 4000 cm⁻¹ the %T (the pen) is at 90%.
8. Now slowly, carefully move the pen carriage manually so that the instrument scans the entire spectrum. Watch the pen! If the baseline creeps up and goes off the paper (Fig. 130), this is not good. Readjust the 100% control to keep the pen on the paper. Now keep going, slowly, and
if the pen drifts up again, readjust it again with the 100% control and get
the pen back on the paper.

9. Now go back to the beginning (4000 cm\(^{-1}\)). If you’ve adjusted the 100% control to get the pen back on the paper at some other part of the spectrum, surprise! The pen will not be at 90% when you get back. This is unimportant. What is important is that the pen stay within the limits, between the 0 and 100%T lines, for the entire spectrum.

10. In any case, if the peaks are too large, with the baseline in the proper place, your sample is just too concentrated. You can wipe some of your liquid sample or mull off one of the salt plates or remake the KBr pellet using less compound or more KBr. Sorry.

11. When you’ve made all the adjustments, press the scan button, and you’re off.

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**Fig. 130** An IR with an unruly baseline.
CALIBRATION OF THE SPECTRUM

Once the run is over, there's one other thing to do. Remember that the index mark on the paper is not exact. You have to calibrate the paper with a standard, usually polystyrene film. Some of the peaks in polystyrene are quite sharp, and many of them are very well characterized. A popular one is an extremely narrow, very sharp spike at 1601.4 cm\(^{-1}\) (6.24 \(\mu m\)).

1. *Don’t move the chart paper* or this calibration will be worthless.
2. Remove your sample, and replace it with the standard polystyrene film sample. You will have to remove any reference beam attenuator and turn the 100% control to set the pen at about 90%, when the chart is at 4000 cm\(^{-1}\).
3. *With the pen off the paper*, move the carriage so that it's just before the calibration peak you want, in this case 6.24 \(\mu m\).
4. Now, quickly, start the scan, *let the pen draw just the tip of the calibration peak*, and quickly stop the scan. You don’t need to draw more than that (Fig. 131). Just make sure you can pick your calibration peak out of the spectral peaks. If it’s too crowded at 1601.4 cm\(^{-1}\), use a different polystyrene peak — 2850.7 or 1028.0 cm\(^{-1}\) (3.51 or 9.73 \(\mu m\)). Anything really well known and fairly sharp will do (Fig. 132).
5. And that’s it. You have a nice spectrum.

IR SPECTRA: THE FINISHING TOUCHES (Fig. 133)

On IR chart paper there are spaces for all sorts of information. It would be nice if you could fill in

2. *Sample*. The name of the compound you’ve just run.
3. *Date*. The day you ran the sample.
4. **Phase.** For KBr, say “solid KBr.” A Nujol mull is “Nujol mull.” Liquids are either solutions in solvents or “neat liquids,” that is, without any solvents, so call them liquids.

5. **Concentration.** For KBr, a solid solution, list milligrams of sample in 100 mg of KBr. For liquids, *neat* is used for liquids without solvents.

6. **Thickness.** Unless you’re using solution cells, *thin film* for *neat liquids*. Leave this blank for KBr samples (unless you’ve measured the thickness of the KBr pellet, which you shouldn’t have done).

7. **Remarks.** Tell where you put your calibration peak, where the sample came from, and anything unusual that someone in another lab might have trouble with when trying to duplicate your work. Don’t put this off until the last day of the semester when you can no longer remember the details. Keep a record of the spectrum *in your notebook*.

You now have a perfect IR, suitable for framing and interpreting.
**Fig. 132** IR of polystyrene film pointing out many calibration peaks.

**Fig. 133** The finishing touches on the IR.
**INTERPRETING IRs**

IR interpretation can be as simple or as complicated as you'd like to make it. You've already seen how to distinguish alcohols from ketones by correlation of the positions and intensities of various peaks in your spectrum with positions listed in IR tables or correlation tables. This is a fairly standard procedure and is probably covered very well in your textbook. The things that are not in your text are

1. *Not forgetting the Nujol peaks.* Mineral oil will give huge absorptions from all the C—H bonds. They'll be the biggest peaks in the spectrum. And every so often, people mistake one of these for something that belongs with the sample.

2. *Nitpicking a spectrum.* Don’t try to interpret every wiggle. There is a lot of information in an IR, but sometimes it is confusing. Think about what it is you’re trying to show, then show it.

3. *Pigheadedness in interpretation.* Usually a case of, “I know what this peak is so don’t confuse me with facts.” Infrared is an extremely powerful technique, but there are limitations. You don’t have to go hog wild over your IR, though. I know of someone who decided that a small peak was an N—H stretch, and the compound had to have nitrogen in it. The facts that the intensity and position of the peak were not quite right, and neither a chemical test nor solubility studies indicated nitrogen, didn’t matter. Oh well.
Nuclear Magnetic Resonance (NMR)
Nuclear magnetic resonance (NMR) can be used like IR to help identify samples. But if you thought the instrumentation for IR was complicated, these NMR instruments are even worse. So I’ll only give some generalities and the directions for the preparation of samples.

For organic lab, traditionally you look only at the signals from protons in your compound, so sometimes this technique is called proton magnetic resonance (PMR). Not naked H+ protons either, eh? The everyday hydrogens in organic compounds are just called protons when you use this technique.

A sample in a special tube is spun between the poles of a strong magnet. A radio-frequency signal, commonly 60 MHz, a little higher than TV Channel 2, is applied to the sample. Now, were all protons in the same environment, there’d be this big absorption of energy in one place in the PMR spectrum. Big deal. But all protons are not the same. If they’re closer to electronegative groups, or on aromatic rings, the signals shift to a different frequency. This change in the position of the PMR signals, which depends on the chemistry of the molecule, is called the chemical shift. Thus, you can tell quite a bit about a compound if you have its NMR.

**LIQUID SAMPLE PREPARATION**

To prepare a liquid sample for NMR analysis,

1. Get an NMR tube. They are about 180 mm long, 5 mm wide, and about a buck apiece for what is euphemistically called the inexpensive model. The tubes are not precision ground, and some may stick in the NMR probe. This should not be your worry, though. They also have matching, color coordinated designer caps (Fig. 134).

2. Get a disposable pipet and a little rubber bulb and construct a narrow medicine dropper. Use this to transfer your sample to the NMR tube. Don’t fill it much higher than about 3–4 cm. Without any solvent, this is called, of course, a neat sample.

3. Ask about an internal standard. Usually tetramethylsilane (TMS) is chosen because most other proton signals from any sample you might have fall at lower frequencies than that of the protons in TMS. Sometimes hexamethyldisiloxane (HMDS) is used because it doesn’t boil
1. Touch lightly to side — let sample go down the side of the tube

2. Fill to at least $\frac{3}{4}$ in.

3. Then cap the tube with the NMR tube cap

Fig. 134 Loading the typical NMR tube.

out of the NMR tube like TMS can. TMS boils at 26–28°C; HMDS boils at 101°C. Add only 1 or 2 drops.

4. Cap the tube and have the NMR of the sample taken. It's really out of place for me to tell you more about NMR here. Buy my next book, *If They Don't Work . . . They're Machines*. 
5. Last point: cleanliness. If there is trash in the sample, get rid of it. Filter it or something, will you?

**SOLID SAMPLES**

Lucky you. You have a solid instead of a liquid. This presents one problem. What are you going to dissolve the solid in? Once it’s a solution, you handle it just like a liquid sample. Unfortunately, if the solvent has protons and you know there’ll be much more solvent than sample, you’ll get a major proton signal from your solvent. Not a good thing, especially if the signals from your solvent and sample overlap.

**Protonless Solvents**

Carbon tetrachloride, a solvent without protons, is a typical protonless solvent. In fact, it’s practically the only example. So if your sample dissolves in CCl₄, you’re golden. Get at least 100 mg of your compound in enough solvent to fill the NMR tube to the proper height.

**CAUTION!** CCl₄ is toxic and potentially carcinogenic. Handle with extreme care.

**Deuterated Solvents**

If your compound does not happen to dissolve in CCl₄, you still have a shot because deuterium atoms do not give PMR signals. This is logical, since they’re not protons. The problem is that deuterated solvents are expensive, so do NOT ask for, say, D₂O or CDCl₃, the deuterated analogs of water and chloroform, unless you’re absolutely sure your compound will dissolve in them. Always use the protonic solvents—H₂O or CHCl₃ here—for the solubility test. There are other deuterated solvents, and they may or may not be available for use. Check with your instructor.
I've included a spectrum of ethylbenzene (Fig. 135) to give you some idea of how to start interpreting NMRs. Obviously, you'll need more than this. See your instructor or any good organic chemistry text for more information.

The Zero Point

Look at the extreme right of the NMR. That single, sharp peak comes from the protons in the internal standard, TMS. This signal is defined as zero, and all other values for the chemical shift are taken from this point. The units are parts per million (ppm), and you use the Greek letter delta (\(\delta\)): \(\delta 0.0\).

Protons of almost all other compounds you’ll see will give signals to the left of zero; positive \(\delta\) values, shifted downfield from TMS. There are compounds that give PMR signals shifted upfield from TMS: negative \(\delta\) values.

Upfield and downfield are directions relative to where you point your finger on the NMR chart.

Signals to the right of where you are are upfield. Signals to the left of where you are are downfield.

The Chemical Shift

You can see that all the peaks don’t fall in the same place, so the protons must be in different surroundings. There is one signal at \(\delta 1.23\), one at \(\delta 2.75\), and another at \(\delta 7.34\). You usually take these values from the center of a split signal (that’s coming up). See that the TMS is really zero before you report the chemical shift. If it is not at zero, you’ll have to add or subtract some correction to all the values. This is the same as using a polystyrene calibration peak to get an accurate fix on IR peaks.

You’ll need a correlation table or a correlation chart (Fig. 136) to help in interpreting your spectrum. The \(-\text{CH}_3\) group is about in the right place (\(\delta 1.23\)). The \(\delta 7.34\) signal is from the aromatic ring and, sure enough, that’s where signals from aromatic rings fall. The \(\delta 2.75\) signal from the \(-\text{CH}_2\) is a bit trickier to interpret. The chart shows a \(-\text{CH}_3\) on a benzene ring in this.
Fig. 135  NMR of ethylbenzene.
**Fig. 136** A garden variety NMR correlation chart.
area. Don’t be literal and argue that you don’t have a $-{1CH}_3$; you have a $-{CH}_2-{CH}_3$. All right, they’re different. But the $-{CH}_2$— group is on a benzene ring and attached to a $-{CH}_3$. That’s why those $-{CH}_2$ protons are further downfield; that’s why you don’t classify them with ordinary R$-{CH}_2$—R protons. Use some sense and judgment.

I’ve blocked out related groups on the correlation table. Look at the set from δ3.1 to δ4.0. They’re the areas that protons on carbons attached to halogens fall in. Read that again. It’s protons on carbons attached to halogens. The more electronegative the halogen on the carbon, the further downfield the chemical shift of those protons. The electronegative halogen draws electrons from the carbon and thus from around the protons on the carbon. These protons, now, don’t have as many electrons surrounding them. They are not as shielded from the big bad magnetic field as they might be. They are de-shielded, so their signal falls downfield.

The hydrogen-bonded protons wander all over the lot. Where you find them, and how sharp their signals are, depends at least on the solvent, the concentration, and the temperature.

Some Anisotropy

So what about aromatic protons (δ6.0–9.5) aldehyde protons (δ9.5–9.6), or even protons on double, nay triple bonds (δ2.5–3.1)? All these protons are attached to carbons with π bonds, double or triple bonds, or aromatic systems. The electrons in these π bonds generate their own little local magnetic field. This local field is not spherically symmetric — it can shield or deshield protons depending on where the protons are — it’s anisotropic. In Fig. 137, the shielding regions have plusses on them, and deshielding regions have minuses.

This is one of the quirks in the numbering system. Physically and psychologically, a minus means less (less shielding), and DOWNfield is further left on the paper; yet the value of δ goes UP. Another system uses the Greek tau (τ) — that’s 10.0–δ. So δ0.0 (ppm) is 10.0τ. Don’t confuse these two systems. And don’t ever confuse deshielding (or shielding) with the proper direction of the chemical shift.
Some NMR Interpretation

Back at ethylbenzene, you’ll find that the —CH₂— and the —CH₃ protons are not single lines. They are split. Spin—spin splitting. Such a fancy name. Protons have a spin of plus or minus ½. If I’m sitting on the methyl group, I can see two protons on the adjacent carbon (—CH₂—). (Adjacent carbon, remember that.) They spin, so they produce a magnetic field. Which way do they spin? That’s the crucial point. Both can spin one way, plus. Both can spin one way, minus. Or, each can go a different way; one plus, one minus.

Over at the methyl group (adjacent carbon, eh?), you can feel these fields. They add a little, they subtract a little, they cancel a little. So your methyl group splits into three peaks! It’s split by the two protons on the adjacent carbon.

Don’t confuse this with the fact that there are three protons on the methyl group! THAT HAS NOTHING TO DO WITH IT! It is mere coincidence.

The methyl group shows up as a triplet because it is split BY TWO protons on the ADJACENT carbon.
Now what about the intensities? Why's the middle peak larger? Get out a marker and draw an A on one proton and a B on the other. OK. There's only one way for A and B to spin in the same direction — *Both A and B are plus* or *both A and B are minus*. But there are two ways for them to spin opposite each other — *A plus with B minus; B plus with A minus*. This condition happens two times. Both A and B plus happen only one time. Both A and B minus happen only one time. So what? So the ratio of the intensities is 1:2:1. Ha! You got it — a **triplet**. Do this whole business sitting on the —CH₂— group. You get a **quartet** — four lines — because the —CH₂— protons are adjacent to a methyl group. They are split BY three to give FOUR lines (Fig. 138).

No, that is not all. You can tell that the —CH₂— protons and the —CH₃ protons split each other by their **coupling constant**, the distance between the split peaks of a single group. Coupling constants are called **J values**, and are usually given in hertz (Hz). You can read them right from the chart, which has a grid calibrated in hertz. If you find protons at different chemical shifts

---

**Fig. 138** Spin alignments for the ethyl group.
and their coupling constants are the same, they’re splitting and coupling with each other.

**Integration**

Have you wondered about those funny curves drawn over the NMR peaks? They’re *electronic integrations* and they can tell you how many protons there are at each chemical shift. Measure the distances between the horizontal lines just before and just after each group. With a cheap plastic ruler I get 52 mm for the benzene ring protons, 21 mm for the —CH₂— protons, and about 30 mm for the —CH₃ protons. Now you divide all the values by the smallest one. Well, 21 mm is the smallest, and without a calculator I get 2.47 : 1 : 1.43. Not even close. And how do you get that 0.47 or 0.43 proton? Try for the simplest *whole number ratio*. Multiply everything by 2, and you’ll have 4.94 : 2 : 2.86. This is very close to 5 : 2 : 3, the actual number of protons in ethylbenzene. Use other whole numbers; the results are not as good and you can’t justify the splitting pattern — 3 split BY 2 and 2 split BY 3 — with other ratios. Don’t use each piece of information in a vacuum.

There are a lot of other things in a typical NMR. There are *spinning sidebands*, small duplicates of stronger peaks, evenly spaced from the parent peak. They fall at *multiples of the spin rate*, here about 30 Hz. Spin the sample tube faster and these sidebands move farther away; slow the tube and they must get closer.

Signals that split each other tend to lean toward each other. It’s really noticeable in the triplet and even distorts the intensity ratio in the quartet some. Ask your instructor or see another textbook if you have questions.
Theory of Distillation
Distillation is one of the more important operations you will perform in the organic chemistry laboratory. It is important that you understand some of the physical principles going on during a distillation. Different systems require different treatments; the explanations that follow parallel the classifications of distillations given earlier in the book.

**CLASS 1: SIMPLE DISTILLATION**

In a simple distillation, you recall, you separate liquids boiling BELOW 150°C at one atmosphere from

1. Nonvolatile impurities.
2. Another liquid boiling at least 25°C higher than the first. The liquids should dissolve in each other. The reason the liquids should dissolve in each other, is that if they do not, then you should treat the system like a steam distillation, and if you’re going to steam distill, be sure to look at the discussion for Class 4: Steam Distillations. The reason the boiling points should have a 25°C difference, is so you may assume the higher boiling component doesn’t do anything but sit there during the distillation. Otherwise, you may have a two-component system, and you also need to look at the discussion for Class 3 Fractional Distillations, as well as here.

That said, let’s go on with our discussion of the distillation of one-component systems.

Suppose you prepared isobutyl alcohol by some means, and at the end of the reaction you wound up with a nice yellow-brown liquid. Checking any handbook, you find that isobutyl alcohol is a colorless liquid that boils at 108.1°C at 760 torr. On an STP day, the atmospheric pressure (P) is 760 torr and the boiling point is the **normal boiling point**. At that point, the single point when the vapor pressure of the liquid is the same as that of the atmosphere, the liquid boils.

Fearing little, you set up a Class 1 Simple Distillation and begin to heat the mix. If you kept track of the temperature of the liquid (and you don’t; the
thermometer bulb is up above the flask), and its vapor pressure, you’d get the temperature-vapor pressure data in columns 1 and 2 of Table 1.

At 82.3°C the vapor pressure of the liquid is 290.43 torr. Much lower than atmospheric pressure. The liquid doesn’t boil. Heat it to, say, 103.4°C and the vapor pressure is 644.16 torr. Close to atmospheric pressure, but no prize. Finally, at 108.1°C we have 760.04 torr. The vapor pressure of the liquid equals that of the atmosphere, and the liquid boils.

Now, do you see an entry in the table for brown gunk? Of course not. The brown gunk must have a very low (or no) vapor pressure at any temperature you might hit during your distillation. Without a vapor pressure, there can be no vapor. No vapor and there’s nothing to condense. Nothing to condense, and there’s no distillation. So the isobutyl alcohol comes over clean and pure, and the brown gunk stays behind.

If you plot the temperature and vapor pressure data given in Table 1, you reconstruct the liquid-vapor equilibrium line in the phase diagram of that liquid (Fig. 139). The equation of this line, and you might remember this from your freshman chemistry course, is the Clausius–Clapyron equation:

\[ p = p^\circ \exp \left\{ \frac{-\Delta H}{R} (1/T - 1/T^\circ) \right\} \]

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Isobutyl alcohol (torr)</th>
<th>Isopropyl alcohol (torr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>82.3</td>
<td>290.43</td>
<td>760.00</td>
</tr>
<tr>
<td>83.2</td>
<td>301.05</td>
<td>786.31</td>
</tr>
<tr>
<td>85.4</td>
<td>328.41</td>
<td>853.90</td>
</tr>
<tr>
<td>86.9</td>
<td>348.27</td>
<td>902.76</td>
</tr>
<tr>
<td>88.7</td>
<td>373.46</td>
<td>964.52</td>
</tr>
<tr>
<td>90.9</td>
<td>406.34</td>
<td>1044.83</td>
</tr>
<tr>
<td>95.8</td>
<td>488.59</td>
<td>1244.30</td>
</tr>
<tr>
<td>99.9</td>
<td>567.96</td>
<td>1435.11</td>
</tr>
<tr>
<td>103.4</td>
<td>644.16</td>
<td>1616.99</td>
</tr>
<tr>
<td>106.2</td>
<td>711.23</td>
<td>1776.15</td>
</tr>
<tr>
<td>108.1</td>
<td>\textbf{760.04}</td>
<td>1891.49</td>
</tr>
</tbody>
</table>
**Fig. 139** Vapor pressure vs. temperature curve for isobutyl alcohol.

**Clausius & Clapyron**

So if you want to know how the vapor pressure of a substance is going to vary with temperature, you can use the *Clausius–Clapyron* equation

\[ p = p^\circ \exp \left\{ \frac{-\Delta H}{R} \left( \frac{1}{T} - \frac{1}{T^\circ} \right) \right\} \]

where

- \( p^\circ \) is a known vapor pressure
- \( T^\circ \) is a known temperature (in K, not °C)
- *These are usually taken from the normal boiling point.*
- \( \Delta H \) is the heat of vaporization of the liquid
- \( R \) is the universal gas constant; (8.314 J/mole-K)
- \( T \) is the temperature you want the vapor pressure for and \( p \) is the vapor pressure, that you calculate, for the temperature you want, \( T \).

I’ve put this formula to two uses:
1. From the normal boiling point of isobutyl alcohol ($T° = 108.1^\circ$C, $381.2$K; $p°=760$ torr), and one other vapor pressure measurement I found while doing research for this section ($T=100^\circ$C, 373K; $p=570$ torr), I've gotten the heat of vaporization ($\Delta H$), the heat needed to vaporize a mole of pure isobutyl alcohol itself. Using these two values in the Clausius–Clapyron equation, the $\Delta H$ for isobutyl alcohol is 10039.70 cal/mole.

2. Now, with this $\Delta H$ for isobutyl alcohol, I've calculated the field of pressures you see in Table 1 from temperatures of 82.3 to 108.1° C. That's why the last pressure at 108.1 is 760.04, and not 760.00. The 760.04 value has been back-calculated using the $\Delta H$ that we calculated from the two pressure-temperature points in the first place.

I've also listed the vapor pressure data for isopropyl alcohol in column 3 of Table 1. Again, two steps were required to generate the data:

1. Two known vapor pressure-temperature points ($T°=82.3$°C, 355.4K, $p°=760$ torr; $T=100$°C, 373K, $p=1440$ torr) were used to calculate the $\Delta H$: 9515.73 cal/mol.

2. Now that $\Delta H$, and temperatures from 82.3 to 108.1° C, were used to calculate a field of vapor pressures for isopropyl alcohol. These are in column 3 of Table 1.

I've done these things for a few good reasons:

1. To show you how to use the Clausius–Clapyron equation, and to show you how well the equation fits over small temperature ranges. The calculated boiling point pressure for isobutyl alcohol (760.04 torr) is not very different from the normal boiling point pressure of 760.00 torr (0.005%).

2. To show you that compounds with higher $\Delta H$'s have lower vapor pressures. This means that it takes more energy to vaporize them.

3. To show you that you can calculate vapor pressures that are above the boiling point of the liquid. They have a slightly different meaning, however. There is no liquid isopropanol at 100° C and 760 torr. The vapor pressure at 100° C is 1440 torr, almost twice the atmospheric pressure. But if we artificially increased the pressure over a sample of isopropyl alcohol (pumped up the flask with compressed air?) to 1440 torr, then
heated the flask, the alcohol would no longer boil at 82.3°C. You’d have to go as high as—did someone say 100°C?—before the vapor pressure of the liquid matched the now pumped-up atmospheric pressure, and the liquid would boil.

4. To show you the theory of the next topic, Class 3, Fractional Distillation.

CLASS 3: FRACTIONAL DISTILLATION

In a fractional distillation, you remember, you are usually separating liquid mixtures, soluble in one another, that boil at less than 25°C from each other at a pressure of one atmosphere.

Now I didn’t discuss both isopropyl and isobutyl alcohol in the last section for my health. Suppose you’re given a mixture of these two to separate. They are miscible in each other and their boiling points are just about 25°C apart. A textbook case, eh?

A Hint from Dalton

So you set up for a fractional distillation and begin to heat the liquid mixture. After a bit, it boils. And what does that mean? The vapor pressure of the solution now (not just one component), is equal to the atmospheric pressure, 760 torr. (We’re very lucky textbook-land has so many STP days.) Each component exerts its own vapor pressure, and when the total pressure reaches 760 torr, the solution boils.

\[ P_{\text{Total}} = P_A + P_B \]

This is Dalton’s Law of partial pressures. The total pressure of the gang is equal to the sum of their individual efforts. Here, A could be the isopropyl alcohol and B the isobutyl, (it doesn’t matter) but \( P_{\text{Total}} \) must be the atmospheric pressure, \( P_{\text{atm}} \). So a special version of Dalton’s Law of partial pressures for use in fractional distillation will be

\[ P_{\text{atm}} = P_A + P_B \]
Dalton and Raoult

If that's all there were to it, we'd be talking about Class 4 Steam Distillations and the like, where the components aren't soluble, and we could quit. Here, the two are soluble in each other. The individual vapor pressures of each component ($P_A$, $P_B$) depend not only on the temperature, but also on their mole fraction.

It makes sense, really. Molecules are the beasties escaping from solution during boiling, and, well, if the two liquids dissolve in each other perfectly, the more molecules (moles) of one component you have, the more the solution behaves like that one component, until it gets to be the same as a one-component liquid. This is Raoult's Law

$$P_A = X_A P_A^o$$

where

- $P_A$ is the vapor pressure of A from the mixture.
- $X_A$ is the mole fraction of liquid A.
- $P_A^o$ is the vapor pressure of the pure liquid A.

If we change the A's to B's, can you still follow me? It's the same thing, only now with liquid B. If we combine the special case of Dalton's Law with Raoult's Law, we get

$$P_{atm} = X_A P_A^o + X_B P_B^o$$

Look at this. If there is NO B, then the fraction of A is 1, and the pure liquid A boils when its vapor pressure equals the atmospheric pressure. Didn't I say that? Similarly, for B without A, the mole fraction of B is 1, and it too boils when its vapor pressure equals the atmospheric pressure.

At this point, you're usually given the temperature versus mole fraction diagram for two miscible liquids (Fig. 140), and you're told it's a consequence of Raoult's Law. Well, yes. But not directly. Raoult's Law is a relationship of pressure, not temperature, versus mole fraction; and Raoult's Law is pretty much a straight line. You don't need all your orbitals filled to see that you've been presented with a temperature versus mole fraction diagram, there are two lines (not one), and neither of them are very straight.
A Little Algebra

I want to convert the combined laws of Dalton and Raoult such that I can show the variation in mole fraction explicitly. First, you’ll agree that the mole fractions of A and B must add to 1 (or they wouldn’t be fractions, eh?), so

\[ X_A + X_B = 1 \]

Now back with the Dalton and Raoult

\[ P_{atm} = X_A P_A^o + X_B P_B^o \]

and seeing that \( X_B = 1 - X_A \), I substitute to get

\[ P_{atm} = X_A P_A^o + (1 - X_A) P_B^o \]

Expand this expression with a multiplication:

\[ P_{atm} = X_A P_A^o + P_B^o - X_A P_B^o \]

Collect the terms with mole fraction in them:

\[ P_{atm} = X_A P_A^o - X_A P_B^o + P_B^o \]

And factor the mole fraction out to give:

\[ P_{atm} = X_A (P_A^o - P_B^o) + P_B^o \]

To isolate the mole fraction \( X_A \) subtract \( P_B^o \) from both sides:

\[ P_{atm} - P_B^o = X_A (P_A^o - P_B^o) \]

and divide by \((P_A^o - P_B^o)\) to get

\[ X_A = \frac{P_{atm} - P_B^o}{P_A^o - P_B^o} \]

Clausius and Clapyron Meet Dalton and Raoult

We still have a formula that relates mole fraction to pressure. Note, however, that with the exception of the atmospheric pressure \((P_{atm})\) all the other pressures are that of pure liquids \((P_A^o \text{ and } P_B^o)\). Now, how does the vapor
pressure of a pure liquid vary with temperature? Smite your forehead and say
that you could have had a V-8. The Clausius–Clapyron equation:

\[ p = p^* \exp \left\{ \frac{-\Delta H}{R} \left( \frac{1}{T} - \frac{1}{T^*} \right) \right\} \]

The \( P^* \)’s of Dalton–Raoult are vapor pressures taken at fixed temperatures. They are the \( p \)'s in the Clausius–Clapyron equation found with a variation in temperature. Don’t believe me? Pick a vapor pressure and temperature pair from Table 1 for either liquid, and let these be \( p^* \) and \( T^* \) (and don’t forget to use K, not °C). Now what happens when the “unknown” temperature (\( T \)) is the same as \( T^* \)? The \((1/T - 1/T^*)\) becomes zero, the entire exponent becomes zero, \( p^* \) is multiplied by 1 (anything to the power zero is 1, eh?) and so \( p = p^* \).

The next part is messy, but somebody’s got to do it. I’m going to use the vapor pressure–temperature data for the normal boiling points of both liquids in the Clausius–Clapyron equation. Why? They’re convenient, known vapor pressure–temperature points. When I do this, though, I exercise my right to use different superscripts to impress upon you that these points are the normal boiling points. So for liquid A, we have \( p_A^* \) and \( T_A^* \); if A is isobutyl alcohol, \( p_A^* = 760 \) torr and \( T_A^* = 101.8 \)°C. For liquid B, we have \( p_B^* \) and \( T_B^* \); if B is isopropyl alcohol, \( p_B^* = 760 \) torr and \( T_B^* = 82.3 \)°C.

Using the new letters, above, and substituting the Clausius–Clapyron equation for every \( P^* \) you get

\[ X_A = \frac{P_{atm} - p_B^* \exp \left\{ \frac{-\Delta H}{R} \left( \frac{1}{T} - \frac{1}{T_B^*} \right) \right\}}{p_A^* \exp \left\{ \frac{-\Delta H}{R} \left( \frac{1}{T} - \frac{1}{T_A^*} \right) \right\} - p_B^* \exp \left\{ \frac{-\Delta H}{R} \left( \frac{1}{T} - \frac{1}{T_B^*} \right) \right\}} \]

Phew!

The only variables in this beast are the mole fraction \((X)\), and the temperature \((T)\). Every other symbol is a constant. We finally have the equation for the bottom line of the temperature-mole fraction diagram, something that has eluded us for years.
Dalton Again

What about the upper curve? Glad you asked (sigh.). The composition in the vapor is also related to Dalton’s Law of Partial Pressures. For an ideal gas

\[ PV = nRT \] (and you thought you’d never see that again!)

and for the vapors above the liquid,

\[ P_A = n_A \left( \frac{RT}{V} \right) \quad \text{and} \quad P_B = n_B \left( \frac{RT}{V} \right) \]

Yet we know that the total pressure of Dalton’s gang is the sum of their individual efforts:

\[ P_{\text{Total}} = P_A + P_B \]

So,

\[ P_{\text{Total}} = n_{\text{Total}} \left( \frac{RT}{V} \right) \]

Now watch, as I divide the pressure of A by the total pressure:

\[ \frac{P_A}{P_{\text{Total}}} = \frac{n_A}{n_{\text{Total}}} \left( \frac{RT}{V} \right) \]

Well the RT/V’s cancel giving

\[ \frac{P_A}{P_{\text{Total}}} = \frac{n_A}{n_{\text{Total}}} \]

The ratio of the number of moles of A to the total number of moles is the mole fraction of component A in the vapor

\[ X_A^{\text{vapor}} = \frac{P_A}{P_{\text{Total}}} \]

\( P_{\text{Total}} \) for the ordinary distillation is the atmospheric pressure, 760 torr (\( P^\star \)'s, eh?). \( P_A \) is the vapor pressure of A, and again by Raoult’s Law, \( P_A = X_A P_A^\star \). Putting the two together, we get

\[ X_A^{\text{vapor}} = X_A^{\text{liquid}} P_A^\star / 760 \]
We can make the same kind of substitution of Clausius–Clapyron here, and get a similarly curved function for the upper line in the temperature-mole fraction diagram.

To show you that all this really does work, I’ve listed the experimental composition data for the isopropyl/isobutyl alcohol system from Landolt–Bornstein (Landolt–Bornstein is to physical chemistry what Beilstein is to organic. And wouldn’t that make for a wild analogy question on the college board entrance exams?), along with my calculated data (Table 2) (That explains my choice of temperatures for Table 1.). I’ve also given the absolute and percent differences between the experimental data, and what I’ve calculated. These differences are on the order of 1% or less, a very good agreement, indeed.

So now, for any two liquids, if you have their normal boiling points and vapor pressures at any other temperature, you can generate the temperature-mole fraction diagram.

**What Does It All Mean?**

Getting back to the temperature-mole fraction diagram (Figure 140), suppose you start with a mixture such that the mole fractions are as follows: isobutyl alcohol, 0.60, and isopropyl alcohol, 0.40. On the diagram, that composition is point A at a room temperature of 20°C. Now you heat the mixture and you travel upwards from point A to point B; the liquid has the same composition, it’s just hotter.

At 95°C, point B, the mixture boils. Vapor, with the composition at point C comes flying out of the liquid (the horizontal line tying the composition of the vapor to the composition of the liquid is the liquid–vapor tie line.), and this vapor condenses (Point C to point D), say, part of the way up your distilling column.

Look at the composition of this new liquid (Point E). It is richer in the lower-boiling component. The step cycle B-C-D represents one distillation.

If you heat this new liquid that’s richer in isopropyl alcohol (Point D), you get vapor (composition at Point G along a horizontal tie-line) that condenses to liquid H. So step cycle D-G-H is another distillation. The two steps represent two distillations.
Table 2  Experimental and Calculated Data.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Normal BP (°C)</th>
<th>Vapor Pressure @ 100 (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-Propanol</td>
<td>82.3</td>
<td>1440 torr</td>
</tr>
<tr>
<td>2-Methyl-1-propanol</td>
<td>108.2</td>
<td>570 torr</td>
</tr>
</tbody>
</table>

Data from Moore (3rd. ed.)

Calculated Δ H (VAP)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Δ H (VAP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-Propanol</td>
<td>9515.73 cal/mol</td>
</tr>
<tr>
<td>2-Methyl-1-propanol</td>
<td>10039.70 cal/mol</td>
</tr>
</tbody>
</table>

Comparison of Data from Moore and Calculated Data (T=100 (°C))

<table>
<thead>
<tr>
<th>Vapor pressures:</th>
<th>Moore</th>
<th>Calc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(torr)</td>
<td>2-Propanol</td>
<td>1440</td>
</tr>
<tr>
<td></td>
<td>2-Methyl-1-propanol</td>
<td>570</td>
</tr>
<tr>
<td>Mole fraction:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(in liquid)</td>
<td>2-Propanol</td>
<td>0.219</td>
</tr>
<tr>
<td></td>
<td>2-Methyl-1-propanol</td>
<td>0.781</td>
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<tr>
<td>Mole fraction:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(in vapor)</td>
<td>2-Propanol</td>
<td>0.415</td>
</tr>
<tr>
<td></td>
<td>2-Methyl-1-propanol</td>
<td>0.585</td>
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</table>

Mole Fraction of 2-Propanol Liquid Data

<table>
<thead>
<tr>
<th>#</th>
<th>T(°C)</th>
<th>X Calc</th>
<th>X Lit.</th>
<th>Diff</th>
<th>% Diff</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>83.2</td>
<td>0.9458</td>
<td>0.9485</td>
<td>-0.0027</td>
<td>-0.286</td>
</tr>
<tr>
<td>2</td>
<td>85.4</td>
<td>0.8213</td>
<td>0.8275</td>
<td>-0.0062</td>
<td>-0.748</td>
</tr>
<tr>
<td>3</td>
<td>86.9</td>
<td>0.7425</td>
<td>0.7450</td>
<td>-0.0025</td>
<td>-0.331</td>
</tr>
<tr>
<td>4</td>
<td>88.7</td>
<td>0.6540</td>
<td>0.6380</td>
<td>0.0160</td>
<td>2.505</td>
</tr>
<tr>
<td>5</td>
<td>90.9</td>
<td>0.5539</td>
<td>0.5455</td>
<td>0.0084</td>
<td>1.540</td>
</tr>
<tr>
<td>6</td>
<td>95.8</td>
<td>0.3591</td>
<td>0.3455</td>
<td>0.0136</td>
<td>3.949</td>
</tr>
<tr>
<td>7</td>
<td>99.9</td>
<td>0.2215</td>
<td>0.2185</td>
<td>0.0030</td>
<td>1.357</td>
</tr>
<tr>
<td>8</td>
<td>103.4</td>
<td>0.1191</td>
<td>0.1155</td>
<td>0.0036</td>
<td>3.096</td>
</tr>
<tr>
<td>9</td>
<td>106.2</td>
<td>0.0458</td>
<td>0.0465</td>
<td>-0.0007</td>
<td>-1.507</td>
</tr>
</tbody>
</table>

Mole Fraction of 2-Propanol Vapor Data

<table>
<thead>
<tr>
<th>#</th>
<th>T(°C)</th>
<th>X Calc</th>
<th>X Lit.</th>
<th>Diff</th>
<th>% Diff</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>83.2</td>
<td>0.9785</td>
<td>0.9805</td>
<td>-0.0020</td>
<td>-0.201</td>
</tr>
<tr>
<td>2</td>
<td>85.4</td>
<td>0.9228</td>
<td>0.9295</td>
<td>-0.0067</td>
<td>-0.723</td>
</tr>
<tr>
<td>3</td>
<td>86.9</td>
<td>0.8820</td>
<td>0.8875</td>
<td>-0.0055</td>
<td>-0.618</td>
</tr>
<tr>
<td>4</td>
<td>88.7</td>
<td>0.8300</td>
<td>0.8245</td>
<td>0.0055</td>
<td>0.663</td>
</tr>
<tr>
<td>5</td>
<td>90.9</td>
<td>0.7615</td>
<td>0.7580</td>
<td>0.0035</td>
<td>0.460</td>
</tr>
<tr>
<td>6</td>
<td>95.8</td>
<td>0.5880</td>
<td>0.5845</td>
<td>0.0035</td>
<td>0.600</td>
</tr>
<tr>
<td>7</td>
<td>99.9</td>
<td>0.4182</td>
<td>0.4270</td>
<td>-0.0088</td>
<td>-2.063</td>
</tr>
<tr>
<td>8</td>
<td>103.4</td>
<td>0.2533</td>
<td>0.2510</td>
<td>0.0023</td>
<td>0.936</td>
</tr>
<tr>
<td>9</td>
<td>106.2</td>
<td>0.1070</td>
<td>0.1120</td>
<td>-0.0050</td>
<td>-4.433</td>
</tr>
</tbody>
</table>
Much of this work was carried out using a special distilling column called a **bubble-plate column** (Fig. 141). Each *plate* really does act like a distilling flask with a very efficient column, and one distillation is really carried out on one physical plate. To calculate the number of plates (separation steps, or distillations) for a bubble-plate column, you just count them!

Unfortunately, the fractionating column you usually get is not a bubble-plate type. You have an open tube that you fill with column packing (see “Class 3: Fractional Distillation”) and *no plates*. The distillations up this type of column are not discreet, and the question of where one plate begins and another ends is meaningless. Yet, if you use this type of column, you do get a better separation than if you used no column at all. It’s *as if* you had a column with some bubble-plates. And if your distilling column separates a mixture as well as a bubble-plate column with two real plates, you must have a column with two *theoretical plates*.

You can calculate the number of theoretical plates in your column if you distill a two-component liquid mixture of known composition (isobutyl and isopropyl alcohols perhaps?), and collect a few drops of the liquid condensed from the vapor at the top of the column. You need to determine the composition of that condensed vapor (usually from a calibration curve of known
compositions versus their refractive indices [see Chapter 22, "Refractometry"], and you must have the temperature-mole fraction diagram (Fig. 140).

Suppose you fractionated that liquid of composition A, collected a few drops of the condensed vapor at the top of the column, analyzed it by taking its refractive index, and found that this liquid had a composition corresponding to point J on our diagram. You would follow the same path as before (B-C-D, one distillation; D-G-H, another distillation) and find that composition J falls a bit short of the full cycle for distillation #2.

Well, all you can do is estimate that it falls at, say, a little more than half of the way along this second tie-line, eh (Point K)? OK then. This column has been officially declared to have 1.6 theoretical plates. Can you have tenths of plates? Not with a bubble-plate column, but certainly with any column that does not have discrete separation stages.

Now you have a column with one-point-six theoretical plates. "Is that good?" you ask. "Relative to what," I say. If that column is six feet high, that's terrible. The Height Equivalent to a Theoretical Plate (HETP) is 3.7 feet/plate. Suppose another column also had 1.6 theoretical plates, but was only 6 inches (0.5 ft) high. The HETP for this column is 3.7 in/plate, and if it were 6 feet high, it would have 19 plates. The smaller the HETP, the more efficient the column is. There are more plates for the same length.

One last thing. On the temperature-mole fraction diagram, there's a point F I haven't bothered about. F is the grade you'll get when you extend the A-B
line up to cut into the upper curve and you then try to do anything with this point. I've found an amazing tendency for some folk to extend that line to point F. Why? Up the temperature from A to B and the sample boils. *When the sample boils, the temperature stops going up.* Heat going into the distillation is being used to vaporize the liquid (heat of vaporization, eh?) and all you get is a vapor, enriched in the lower-boiling component, with the composition found at the end of a horizontal tie-line.

**Reality Intrudes I: Changing Composition**

To get the number of theoretical plates, we fractionally distilled a known mixture and took off a small amount for analysis, so as not to disturb things very much. You, however, have to fractionally distill a mixture and hand in a good amount of product, and do it within the time limits of the laboratory.

So when you fractionally distill a liquid, you *continuously* remove the lower boiling fraction from the *top* of the column. And where did that liquid come from? The boiling liquid at the *bottom* of the column. Now if the distillate is richer in the lower boiling component, what happened to the composition of the boiling liquid? I'd better hear you say that the boiling liquid gets richer in the *higher-boiling* component (Fig. 142).

*Fig. 142* Changing composition as the distillation goes on.
So as you fractionally distill, not only does your boiling liquid get richer in the higher-boiling component, so also does your distillate, your condensed vapor. Don’t worry too much about this effect. It happens as long as you have to collect a product for evaluation. Let your thermometer be your guide, and keep the temperature spread less than 2°C.

**Reality Intrudes II: Nonequilibrium Conditions**

Not only were we forced to remove a small amount of liquid to accurately determine the efficiency of our column, we had to do it very slowly. This allowed the distillation to remain at equilibrium. The throughput, the rate at which we took material out of the column, was very low. Of all the molecules of vapor that condensed at the top of the column, most fell back down the column; few were removed. A very high reflux ratio. With an infinite reflux ratio (no throughput), the condensed vapor at the top of the column is as rich in the lower-boiling component as it’s ever likely to get in your setup. As you remove this condensed vapor, the equilibrium is upset, as more molecules rush in to take the place of the missing. The faster you distill, the less time there is for equilibrium to be reestablished — less time for the more volatile components to sort themselves out and move to the top of the column. So you begin to remove higher-boiling fractions as well, and you cannot get as clean a separation. In the limit, you could remove condensed vapor so quickly that you shouldn’t have even bothered using a column.

**Reality Intrudes III: Azeotropes**

Occasionally, you’ll run across liquid mixtures that cannot be separated by fractional distillation. That’s because the composition of the vapor coming off the liquid is the same as the liquid itself. You have an azeotrope, a liquid mixture with a constant boiling point.

Go back to the temperature-mole fraction diagram for the isopropyl alcohol–isobutyl alcohol system (Fig. 140). The composition of the vapor is always different from that of the liquid, and we can separate the two compounds. If the composition of the vapor is the same as that of the liquid, that separation is hopeless. Since we’ve used the notions of an ideal gas in deriving
our equations for the liquid and vapor compositions (Clausius–Clapyron, Dalton, and Raoult), this azeotropic behavior is said to result from deviation from ideality, specifically deviations from Raoult’s Law. Although you might invoke certain interactive forces in explaining nonideal behavior, you cannot predict azeotrope formation a priori. Very similar materials form azeotropes (ethanol–water). Very different materials form azeotropes (toluene–water). And they can be either minimum-boiling azeotropes or maximum-boiling azeotropes.

**Minimum-Boiling Azeotropes**

The ethanol–water aze trope (95% ethanol–5% water) is an example of a minimum boiling azeotrope. Its boiling point is lower than that of the components (Fig. 143). If you’ve ever fermented anything and distilled the results in the hopes of obtaining 200 proof (100%) white lightning, you’d have to content yourself with getting the azeotropic 190 proof mixture, instead. Fermentation usually stops when the yeast die in their own 15% ethanol solution. At room temperature, this is point A on our phase diagram. When you heat the

![Fig. 143 Minimum-boiling ethanol–water azeotrope.](image-url)
solution, you move from point A to point B and, urges to go to point F notwithstanding, you cycle through distillation cycles B-C-D and D-E-G, and, well, guess what comes off the liquid? Yep, the azeotrope. As the azeotrope comes over, the composition of the boiling liquid moves to the right (it gets richer in water), and finally there isn’t enough ethanol to support the azeotropic composition. At that point, you’re just distilling water. The process is mirrored if you start with a liquid that is >95% ethanol and water. The azeotrope comes off first.

Maximum-Boiling Azeotropes
The chloroform–actone azeotrope (52%chloroform–48%acetone) is an example of the much rarer maximum boiling azeotrope. It’s boiling point is higher than that of the components (Fig. 144). At compositions off the azeotrope, you do distillations A-B-C or D-E-F (and so on) until the boiling liquid composition reaches the azeotropic composition. Then that’s all that comes over. So, initially, one of the components comes off first.

Azeotropes on Purpose
You might think the formation of azeotropes to be an unalloyed nuisance, but they can be useful. Toluene and water form a minimum-boiling azeotrope (20.2%water; 85°C). If you needed very dry toluene for some reason, all you need to do is distill some of it. The water–toluene azeotrope comes off first,
and, well, there goes all the water. It gets removed by azeotropic distillation. The technique can also be used in certain reactions, including the preparation of amides from very high boiling acids and amines (they can even be solids.). You dissolve the reagents in toluene, set up a reflux condenser fitted with a Dean–Stark trap, (Fig. 145) and let the mixture reflux. As the amide forms, water is released and the water is constantly removed by azeotropic distillation with the toluene. The azeotrope cools, condenses, and collects in the Dean–Stark trap. At room temperature, the azeotrope is said to “break,” and the water forms a layer at the bottom of the trap. Measure the amount of water and you have an idea of the extent of the reaction.

Absolute (100%) ethanol is often made by adding benzene to the ethanol–water binary azeotrope (two components), to make a ternary azeotrope (three components). This ternary alcohol–water–benzene (18.5:7.4:74.1) azeotrope comes over until all the water is gone, followed by a benzene–ethanol mixture. Finally, absolute ethanol gets its chance to appear, marred only slightly by traces of benzene.

Other Deviations

The furfural–cyclohexane phase diagram (Fig. 146) shows that you can have mixtures that exhibit nonideal behavior, without having to form an azeotrope. In sum, without the phase diagram in front of you, you shouldn’t take the distillation behavior of any liquid mixture for granted.

CLASS 4: STEAM DISTILLATION

Steam distillation is for the separation of mixtures of tars and oils, and they must not dissolve much in water. If you think about it a bit, this could be considered a fractional distillation of a binary mixture with an extreme deviation from Raoult’s Law. The water and the organic oils want nothing to do with each other. So much so, that you can consider them unmixed, in separate compartments of the distilling flask. As such, they act completely indepen-
Fig. 145 Removing water by azeotropic distillation.
Fig. 146 Other deviant behavior (but no azeotropes) in the furfural–cyclohexane system.

dently of each other. The mole fraction of each component in its own compartment is 1. So Raoult’s Law becomes

\[ P_{\text{Total}} = P_A + P_B \]

This is just Dalton’s Law of partial pressures. \( P_{\text{Total}} \) is \( P_{\text{atm}} \) for a steam distillation. So the vapor pressure of the organic oil is now less than that of the atmosphere and the water, and codistills at a much lower temperature.

As an example, suppose you were to try to directly distill quinoline. Quinoline has a boiling point of 237°C at 1 atm. Heating organic molecules to these temperatures may often be a way to decompose them. Fortunately, quinoline is insoluble in water and it does have some vapor pressure at about the boiling point of water (10 torr at 99.6°C). If it had a much lower vapor pressure at the boiling point of water, (say 0.1 torr), there wouldn’t be enough of it vaporizing to make even steam distillation worthwhile.

Well, at 99.6°C, quinoline contributes 10 torr to the total vapor pressure, and water must make up the difference (750 torr) in order to satisfy Dalton’s Law of partial pressures to make \( P_{\text{Total}} \) 760 torr at boiling. Using the relationships for the composition of the vapor over a liquid, we can calculate the quinoline/water ratio coming over.
If we consider each to be an ideal gas, then

\[ PV = nRT \text{ (yes, again)} \]

The number of moles \( n \) of anything is just the weight in grams \( g \) divided by the *molecular weight* of that substance \( (MW) \), and so

\[ PV = \frac{g}{MW}(RT) \]

Multiplication by \( MW \) gives

\[ (MW)PV = gRT \]

and isolating the mass of the material by dividing by \( RT \) gives

\[ (MW)PV/RT = g \]

Now for two vapors, A and B, I’ll construct a ratio where

\[ \frac{(MW)_A P_A V/RT = g_A}{(MW)_B P_B V/RT = g_B} \]

With A and B in the same flask, \( R, T, \) and \( V \) must be the same for each, and we can cancel these terms giving

\[ \frac{(MW)_A P_A}{(MW)_B P_B} = \frac{g_A}{g_B} \]

If we plug in values for the molecular weights and vapor pressures of quinoline and water, we get

\[ \frac{(129)(10)}{(18)(750)} = 0.0956 \]

So, as an approximation, for every 10 g of distillate we collect, 1 g will be our steam distilled quinoline.
Several small extractions are better than one big one." Doubtless you've heard this many times, but now I'm going to try to show that it is true.

By way of example, let's say you have an aqueous solution of oxalic acid, and you need to isolate it from the water by doing an extraction. In your handbook, you find some solubilities of oxalic acid as follows: 9.5g/100g in water; 23.7g/100g in ethanol; 16.9g/100g in diethyl ether. Based upon the solubilities, you decide to extract into ethanol from water, forgetting for the moment that ethanol is soluble in water, and you must have two insoluble liquids to carry out an extraction. Chagrined, you choose diethyl ether.

From the preceding solubility data we can calculate the distribution, or partition coefficient for oxalic acid in the water–ether extraction. This coefficient (number) is just the ratio of solubilities of the compound you wish to extract in the two layers. Here,

$$K_p = \frac{\text{solubility of oxalic acid in ether}}{\text{solubility of oxalic acid in water}}$$

which amounts to 16.9/9.5, or 1.779.

Imagine you have 40 g of oxalic acid in 1000 ml of water, and you put that in contact with 1000 ml of ether. The oxalic acid distributes itself between the two layers. How much is left in each layer? Well if we let $x$ g equal the amount that stays in the water, 1.779$x$ g of the acid has to walk over to the ether. And so

$$\text{Wt of oxalic acid in ether} = (1000\text{ml})(1.779x \text{ g/ml}) = 1779x \text{ g}$$

$$\text{Wt of oxalic acid in water} = (1000\text{ml})(x \text{ g/ml}) = 1000x \text{ g}$$

The total weight of the acid is 40 g (now partitioned between two layers) and

$$2779x \text{ g} = 40g$$

$$x = 0.0144$$

and

$$\text{Wt of oxalic acid in ether} = 1779 (0.0144)\text{g} = 25.6\text{g}$$

$$\text{Wt of oxalic acid in water} = 1000 (0.0144)\text{g} = 14.4\text{g}$$

Now, let's start with the same 40 g of oxalic acid in 1000 ml of water, but this time we will do three extractions with 300 ml of ether. The first 300 ml portion hits, and

$$\text{Wt of oxalic acid in ether} = (300\text{ml})(1.779x \text{ g/ml}) = 533.7x \text{ g}$$
Wt of oxalic acid in water = \( (1000 \text{ml})(x \text{ g/ml}) = 1000x \text{ g} \)

The total weight of the acid is 40 g (now partitioned between two layers) and

\[
1533.7x \text{ g} = 40g \\
x = 0.0261
\]

and

Wt of oxalic acid in ether = 533.7 (0.0261)g = 13.9g

Wt of oxalic acid in water = 1000 (0.0261)g = 26.1g

That ether layer is removed, and the second jolt of 300 ml fresh ether hits, and

\[
Wt \text{ of oxalic acid in ether} = (300\text{ml})(1.779x \text{ g/ml}) = 533.7x \text{ g} \\
Wt \text{ of oxalic acid in water} = (1000\text{ml})(x \text{ g/ml}) = 1000x \text{ g}
\]

But here, we started with 26.1 g of the acid in water, (now partitioned between two layers) and

\[
1533.7x \text{ g} = 26.1g \\
x = 0.0170
\]

and

Wt of oxalic acid in ether = 533.7 (0.0170)g = 9.1g

Wt of oxalic acid in water = 1000 (0.0170)g = 17.0g

Again, that ether layer is removed, and the third jolt of 300 ml fresh ether hits, and

\[
Wt \text{ of oxalic acid in ether} = (300\text{ml})(1.779x \text{ g/ml}) = 533.7x \text{ g} \\
Wt \text{ of oxalic acid in water} = (1000\text{ml})(x \text{ g/ml}) = 1000x \text{ g}
\]

But here, we started with 17.0 g of the acid in water, (now partitioned between two layers) and

\[
1533.7x \text{ g} = 17.0g \\
x = 0.011
\]

and
Wt of oxalic acid in ether = 533.7 (0.011)g = 5.87g

Wt of oxalic acid in water = 1000 (0.011)g = 11.0g

(They don’t quite add up to 17.0g—I’ve rounded them off a bit.)

Let’s consolidate what we have. First, 13.9 g, then 8.5 g and, finally 5.34 g of oxalic acid, for a total of 28.9 g of acid, extracted into 900 ml of ether. OK, that’s not far from 24.7 g extracted once into 1000 ml of ether. That’s because the distribution coefficient is fairly low. But it is more. That’s because several small extractions are better than one large one.
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