Modus Operandi (1-4)

1. TA responsibilities.

**General:** For safety, TAs permit a maximum of fourteen students per section and they require students to wear lab goggles and closed-toed-shoes. TAs ensure that students finish their experiments in a timely fashion (4 hrs max!).

**Instruction and score:** In each experiment class, TAs give a brief overview of the theory and procedure for the daily activities, give a quiz, grade the pre-lab, and in-lab parts, and assign a technique grade. After each experiment, TAs will request a photocopy of the entire experiment (pre-lab, in-lab, and post-lab write-up) for grading. TAs should complete their grading within one week after receiving each lab report. During every experiment, TAs assign 2-3 low technique scores (50) and 2-3 outstanding technique scores (90) to students. TAs use an identical 6A grading spread sheet to track their grades.

**Absence and make-up:** TAs oversee all nine experiments. TAs do not have the authority to cancel class or permit a student to skip a lab. If a student can't attend, TAs have to inform to the instructor with a reasonable reason, who may provide a ‘permission’ email. If space is available, TAs in other sections will contact the student to inform them of potential time to make-up the experiment. Make-up will be allowed up to 2 times in entire course. If a TA is sick, or must miss a lab, they will make arrangements to cover the sections and inform the instructor.

2. General procedures for students in each lab. Each class usually begins with a 5-10 minute quiz that emphasizes important aspects of the current experiment or specifics from the past lab. The TAs will give a 10-30 minute overview of the experiment, including the theory behind the practice; make notes in your notebook. While you are setting up your experiment, the TA will grade the pre-lab of your notebook in red ink, assigning a score with the date and their initials on the first page of the experiment. Before leaving lab, your TA will again initial in-lab of your notebook to verify your experimental procedure and observations. You should work independently on ALL the experiments and write-ups.

Course Outline

Exp. 1 (esterification - laurate)
Exp. 2 (oxidation - cyclohexanone)
Exp. 3 (ketone olefination - stilbene)
Exp. 4 (hydroboration - octanol)
Exp. 5 (Grignard 1,2-add. - triphenylmethanol)
Exp. 6 (Diels-Alder - cyclohexene)
Exp. 7 (Br₂, dehydrohalogenation - diphenylacetylene)
Exp. 8 (condensation - benzoin)
Exp. 9 (oxidation - benzil)
Exp. 10 (Aldol, Diels-Alder - hexaphenylbenzene)
Exp. 11 (amide acylation, Br₂ - 4-bromocotaniilide)
Exp. 12 (glucose acetylations - glucose penta-acetates)
Exp. 13 (diazonium coupling - methyl orange and red)
Exp. 14 (Friedel-Crafts - 2-(4-toluoyl)-benzoic acid)
Exp. 15 (Friedel-Crafts - 2-methylantraquinone)
3. Lab Reports.

**General:** The laboratory notebook is a record of all work performed in the laboratory. It is a legal document that gives testimony of performed work. It should be concise, written clearly and neatly, so as to allow a future experimentalist to reproduce the experiment. Your ability to keep a good notebook can affect also your technique grades.

**Notebook:** Students should use a non-perforated notebook. Do not use a spiral/loose-leaf notebook, and never remove pages from your notebook. The notebook should begin with a table of contents (see page 3 marginal note), including the title of every experiment and the corresponding page number, listed in sequential order. All entries should be written in ink, whether done in advance of the experiment or while making observations. You should never type reports. All pages are to be numbered sequentially. The lab record (pre-, in- and post-labs) is written on only right hand pages of the notebook. This leaves the left-hand pages for notes and extraneous calculations.

**Contents:** Reports consist of three components. The pre-lab (Part I) is completed before coming to lab. During lab, TA lecture notes are then written into the notebook (left-side) and the actual in-lab procedure (Part 2a) is completed on the right-side. The post-lab experimental analysis (Part 2b) is completed after the experiment is done. Part I will be initialed, dated and graded by the TA at the beginning of each experiment. Part I includes: A) a title, the date, your name, you perm number, and your TAs name; B) a purpose/objective; C) a reaction diagram or flow chart; D) table of reagents and products; and E) a brief intended procedure. Throughout the experiment, annotations and changes can be made to Part I and Part 2a on the left hand page in ink. Part 2a is written during the lab. It is comprised of A) the actual experimental procedure and B) experimental observations. Before leaving lab you must get your TAs initials to verify you have composed an experimental procedure and have maintained experimental observations during lab. Part 2b, the final post lab write-up, is written after lab. It includes A) your results, B) a discussion of your results, C) a conclusion, and D) references for any literature used. Any

### Grading

Notebooks (100pts) [25%]  
Part 1 (30pts)  
Part 2a (20pts)  
Part 2b (50pts)  
Technique [10%]  
(50, 70 or 90/100 pts)  
Quizzes [55%]  
Products [10%]  
(60–100/100 pts)

This class is curved. The average for each section of chem 6b is set to a [B]. A few students (1-5) will get A’s and a few students (1-5) will get C’s. The instructor in-charge assigns final grades after evaluating the notebooks, TA grade sheets and in consultation with each TA.
loose paper (spectra and quizzes) and TLC should be neatly taped into the notebook following to the appropriate lab.

**Submission: <lab report for each lab>** A photocopy of Part 1 and Part 2a/b including spectra are submitted to the TA for grading by Friday for the previous Monday and Tuesday labs and by Monday for the previous Wednesday. Thursday and Friday labs. Your TA can reduce your notebook grade by 10% for each day it is late. Experiments not submitted by the beginning of exam week will receive a score of zero.

**<whole lab notebook>** You are required to turn your notebook over to your TA at the end of the quarter. It will be used to assist the instructor in assigning your final letter grade. If a student does not turn in their notebook, they should be prepared to lose at least 2/3 of a letter grade from their final letter grade. At the end of the course please write one page about the course, its strengths and weaknesses, as the last page of your notebook. Make suggestions about what you would you do to improve it.

**Detail of notebook preparation:**
---Pre-Lab [Part 1]--- (30%): need to be done before the beginning of lab
A. Title, date, name, student perm number and TA overseeing the experiment
B. Purpose/Objective
   Give a brief introduction to the purpose of the experiment and the approach to be used. Demonstrate that you understand the objective and the key concepts of the experiment. **Do not copy directly from the laboratory manual.** Usually, one or two paragraphs will be adequate (less than 1/2 a page). Use only the **third person, present tense, passive voice** when writing the introduction.
   (ex) Correct: Cyclohexanol is converted to cyclohexene using......
   Incorrect: In this experiment, I will be performing an acid catalyzed dehydration.... Example: Cyclohexene is prepared from purified cyclohexanol by acid catalyzed dehydration. Trace acid is neutralized with sodium carbonate and the product is salted out of the aqueous extract using brine. The organic layer is dried over magnesium sulfate as a drying reagent. Purified cyclohexene is obtained by simple distillation and it is characterized using IR and NMR spectroscopy by monitoring the loss of -OH and the gain of an alkene.
C. Reaction Diagram
   Relevant, balanced, and fully labeled, chemical equations should be included.
D. Table of Reagents and Products
   Table of reagents needs to be **completed before Lab starts**.
E. Intended Procedure with Flow Chart
   Demonstrate that you are prepared for lab by giving a brief description of what you actually intend to do in lab experimentally. A “game plan” or checklist, written in your own words, will save you time in lab. This can be written in paragraph form or as a bulleted list. Do not copy directly from the laboratory manual. The flow chart is helpful to understand the exact procedure of the lab. Beautiful drawing of the equipments is highly recommended.
   (ex) Weigh lauric acid. Add to 3mL conical vial with stir bar. Add EtOH. Add AcCl. Reflux with water condenser and drying tube for 1 hr...

---In-Lab-[Part 2a]--- (20%)
A. Actual Procedure
   This is an account of what really was done. Do not regurgitate the laboratory manual. Students need to write whole the procedure as exactly carried out. If the procedure has been modified, or changed in any way from the original way written in lab manual, note the changes here. Remember that the procedure section should be sufficiently detailed, such that another student would be able to repeat the whole experiment based on your report. Keep the following points in mind:
   (i) Use the third person, the passive voice, and the **past tense**.
      Correct: The solution was heated on a hot-plate for 30 minutes.
      Incorrect: I heated the solution on a hot-plate for 30 minutes.
      Incorrect: The solution is heated on a hot plate for 30 minutes.
   (ii) Avoid the “recipe format”.
      Incorrect: Heat the solution on a hot-plate for 30 minutes.
   (iii) Incorporate your **observations** into the procedure.
      (ex) The solution was heated on a hot-plate for 30 minutes, during which time the color of the solution changed from red to green.
   (iv) Should be written **concisely** written. Avoid unnecessary detail.
      Correct: 20 mL of hydrochloric acid (3M) was added to the solution with constant stirring.
      Incorrect: 20 mL of 22.5 °C hydrochloric acid (3M) was poured from a graduated cylinder into a 100-mL beaker containing the solution. During this process the solution in the beaker was stirred with a 15-cm long glass rod having a diameter of 5 mm.
B. Observations
   Prepare a simple flow chart of the procedure, and record any observations alongside. This will show your **scientific engagement** and need to be considered as an important matter.
   Correct: The reaction mixture turned green and a precipitate formed. The crude product, a yellow crystal, weight 15mg.

---Post-Lab-[Part 2b]--- (50%)

---Pre-Lab [Part 1]---  (30%): need to be done before the beginning of lab
A. Title, date, name, student perm number and TA overseeing the experiment
A. Results:
This is one of the most important sections of your report. Wherever possible, tabulate your data, such as the melting/bubbling point with its range, any IR and/or NMR spectra, and any other observations or measurements. Include all the spectra, which will be provided from your TA as standards, with your interpretations and peak assignments. Especially, show clearly how did you calculate the % yield.

B. Discussion
This section should be completely based on your results (measured or calculated values) and observations. Whatever the value you got is your data, and you need to consider the meaning of your data. You also need to show your understanding of the experimental mechanistic background. Often you need to site references where you can obtain the supporting informations.
First, your discussion should state what you've made (draw the structure and name it) and what it appears like (was it as expected, compared to a standard or the literature, e.g. white shiny crystalline solid).
Next, discuss the yield and purity of the product(s) you recovered/synthesized. Qualitatively assess the performance. A discussion should quote actual experimental values and not talk in vague terms.
Correct: The product obtained was found to be fairly pure, as it had a mp of 110-112°C, a mp range of only 2°C. This result was 3 degrees below the literature value of 115°C for compound X. This also shows that the product was not completely pure.
Correct: The infrared spectrum of the alkene product (see page xx of this report) had the absorption bands of the expected alkene. 3050 cm⁻¹ sp² C-H stretch and 1650 cm⁻¹ =C=C absorption. No broad alcohol band was observed at 3300 cm⁻¹, indicating no reagent alcohol remains and that the reaction resulted in the conversion of the alcohol to the alkene product. Incorrect: The product obtained was found to be pure. Data interpretation should demonstrate a clear understanding of the technique, the experiment, and the spectra.
The next section of your discussion covers sources of error and loss. Try to think of at least two sources of each. Sources of error include theoretical sources, such as the reaction did not go to 100% completion, and practical sources, such as the instrument or glassware used was not calibrated. Sources of loss include theoretical sources, such as reaction byproduct formation, and practical sources, such as surface adhesion, loss on glassware, and mechanical transfer loss (e.g. spill).
Finally, mention at least one way to improve the experiment.

C. Conclusion
Give your signature and a pledge that all of the observations and conclusion herein are your own and that you believe them to be correct. Then have another person witness your pledge with their signature.

D. References:
You should reference any literature used in your report, i.e. melting points, spectral data, etc. Use an acceptable scientific journal style/format for your references. Be consistent. Author name (surname, initials.), year published. Title, publisher name, publisher location, page numbers

4. Technique Score. During each experiment technique scores will be assigned out of 100 pts; 3-4 low technique scores (50 pts) and 2-3 outstanding technique scores (90 pts) will be assigned to deserving students. The technique score can be very subjective. Your TA can give you a low score if you 1) are wearing inappropriate clothing, 2) have a messy area, 3) use incorrect disposal techniques, 4) are extremely inefficient in lab, 5) have an unorganized unkept notebook, 6) are not wearing your safety goggles, 7) are unprepared for lab. Failure to listen, learn, and comply will be reflected in your technique grade. Your TA can give you an outstanding score if you 1) clean up a dirty area, 2) ask good questions, 3) have a very neat, organized notebook, 4) are efficient during lab. TAs will strive to have everyone finish the lab course with an average technique score. Therefore, if you receive a low score, you should make use of opportunities (preferably in the same lab) to zero it out with a high score.

5. Product Score. Product scores will be assigned out of 60-100 pts for each experiment based on purity and product amount. Purity and product amount will be assessed separately, in relation to other students in the class, on a distribution of 30-50 pts and then the two scores are added together.

6. Grading. Grades are based on in class quizzes, notebooks, technique scores, and product scores. TAs will set the average grade of their section to a 2.9 (B). TAs will enter fifteen notebook scores, ten quiz scores, fifteen product scores, and up to fourteen technique scores. All TAs use the same grading schemes and the same Excel® chem 6B spread sheet. Students are to turn in their notebooks to their TA at the end of the quarter. The grade sheet is emailed to the instructor at the beginning of dead week. No reports or notebooks are accepted after the end of dead week. The TA brings his/her students’ notebooks and meets with the instructor to discuss the final grade of each student.

7. Missing Lab. Missing lab is strongly discouraged. If you miss a lab for an excusable reason, you must contact the instructor (not the TA) within 24 hours to make arrangements for a make-up experiment in another TAs section. If excused, the instructor will
contact other sections and the TA will contact you if they have space. TAs will not give unknown students entry without a permission email from the instructor. You must make-up the old experiment with minimal guidance from the new TA, who may be directing a different experiment. Because of waste disposal issues, if you do not make up the experiment within one week, you will receive “0’s” for Part 1 and Part 2a/b. The new TA must initial and grade Part 1 of your notebook for the experiment that you are performing. The quiz in the other section does not count toward your grade. Your old TA will leave a blank for your missed quiz score. Be sure that your original TA enters scores for Part 1 of the make-up into their grade-book. You’ll submit Part 2a/b to your original TA. However, even if you are allowed to make-up three or more labs, you will probably receive an F as a letter grade.

8. Rules for the Disposal of Reaction Wastes. Disposal of reaction wastes can affect your technique score! Dilute aqueous wastes containing only acids, bases, or salts (which have been neutralized!) may be disposed in sinks. Wet methanol, acetone and ethanol are considered aqueous waste. When mixed with dry organic, however, these solvents are considered organic waste. All organic wastes must be poured into the bottles provided. Please note that their may be separate bottles for halogenated and non-halogenated wastes. Bottles with specific labels for each experiment’s waste will be available in the hood. Solids, such as drying agents, are placed in the plastic bags provided or in solid waste containers. TA’s will independently dispose of your product vials after grading. Do not leave any unlabeled vials with chemicals in your drawer. At the end of the quarter, you must clear all chemicals out of your locker. When in doubt about how to dispose of something, ask your teaching assistant. Also, see the comments concerning waste given at the end of each experiment.

9. Laboratory Safety. Laboratory safety can effect your technique score! Dealing with chemicals requires an alertness and awareness of the problems associated with the handling of volatile, flammable, corrosive and toxic materials. Many generations of organic chemists have learned how to do chemistry both safely and enjoyably. It is necessary to be always cautious, but not to the detriment of performing the experiments expeditiously. Learn to be aware of the safety requirements, but then to enjoy the experience of preparing materials and analyzing them as efficiently as possible.

Syringe disposal. Your TA will give you 1-2 capped needles during the experiments requiring syringes. Upon disposal, 10-20 points will be subtracted from your quiz grade as an “insurance.” When you return the needle, 20-30 points will be re-added to your quiz grade. TA’s will put waste needles in the assigned disposal jars. **DO NOT PUT NEEDLES IN THE TRASH!! REPORT NEEDLES IN THE TRASH TO THE TA.**

Safety glasses. Safety glasses must be worn in the laboratory. You will not be admitted into the laboratory unless your eyes are and remain protected. Visitors must also wear safety glasses. Do not wear contact lenses in the laboratory. Organic fumes may harm them, and caustic reagents cannot be washed from the eye if contact lenses are worn.

Gloves. Gloves should be worn if you are handling corrosive materials. Surgical gloves will not protect you against strong acids, but they are available for your use at other times. Heavy rubber gloves are available for handling extremely corrosive materials. You may wish to purchase your own rubber gloves and keep them in your locker.

Shoes. Sandals or open shoes are forbidden in the laboratory. Clothing which leaves your legs exposed should not be worn, unless a laboratory coat or apron is worn as well.

Hair. Your hair should be pulled and tied back from the face so that it cannot be caught in equipment or open flames.

Glassware. Glassware is cleaned easiest after every use. Most organic materials are removed from glassware with acetone and water. Soap may not be necessary.

Heating. Heat is an ignition source. Only use heat in a working hood. Never heat a closed system! Keep flammable organic solvents away from flames and sources of heat. Ether has a very low flash point and may be ignited by a hot-plate.

Cleanliness. Your locker and bench-top should always be ordered and neat. Do not store chemical or reagents in your locker, with the exception of labeled products and recrystallizations. The balance-area should be cleaned after every use. Clean up any spills immediately. Organic solvents often dissolve plastics and rubber items.

Accidents. Showers and eye washes are available. The eye wash fountains at the front of the lab are for flushing the eyes with water after an accident.
10. Laboratory Instruments

The IR: The Jasco FT-IR is a powerful, but easy to use instrument. First, you will need to acquire background scans by selecting “Background” from the Scan menu. Make sure that nothing is in the sample chamber when this is done. To run a sample, place the plate in the V-shaped sample holder that permanently resides within the FT-IR. Select “Sample Scan” from Scan menu. From the File menu, select “Plot”. In the resulting window, select “Plot” once again. Click “Done” once plotting has begun.

Because of their expense and moisture sensitivity, the single crystal salt plates should be handled carefully by the edges. You will mostly use “the thin film technique” for your dry samples. To do this, set one salt plate flat on a clean surface (e.g. paper towel, Kimwipe, etc.), put one small drop of “neat” (undiluted) sample on the plate, and one drop of methylene chloride to evenly disperse the sample. Let it dry on the plate. You can now run your sample. After recording your spectra, clean the salt plate with dichloromethane and Kimwipes, touching the edges only. Put the plate back into the desiccator, or give it to the next student in line.

KBr pellets are obtained by using the metal hexagonal nut and tightening the bolts with a torque wrench to approximately 30-40 lbs/square inch. The most important part of the preparation of the pellet is to see that your dry sample is ground well (approximately 5 minutes grinding) and that about 10 times as much dry KBr is added to the mortar and mixed well. If everything is dry, when you carefully remove the screw on the die, you will see an almost transparent pellet that may be mounted by placing the die on the plastic cell holder.

The GC: Your TA will show you how to use the syringe, the recorder, and the gas chromatograph for the separation of your products during the elimination experiment.

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Forward to the chem 6b student

Experimental techniques and instrumentation are the backbone of experimental organic chemistry. These experiments expose you to the fundamental techniques used by organic chemists. If you were to venture into a research corridor of the chemistry building, graduate students (your TAs) are taking melting points, performing distillations and extractions, using chromatography, and employing NMR and IR spectroscopy on a regular basis. Their apparatus may be more advanced, but the basic principles remain unchanged. Your TA serves as your well suited guide on this adventure. When you complete this course, we hope that you will further understand these principles. We hope to see you soon in Chem 6c or in one of our research labs soon. Have fun.
Review of spectroscopy

Infra Red.

- IR is good for distinguishing FGI
- An infrared spectrum is obtained by passing infrared radiation through the sample
  - Wavenumber ($\nu$) is another way to describe the frequency of electromagnetic radiation.
  - High frequencies, large wavenumbers ($\nu$), and short wavelengths are associated with high energy
  - The covalent bonds in molecules are constantly vibrating and each stretching and bending vibration of a bond occurs with a characteristic frequency
  - The greater the change in dipole moment, the more intense the absorption.
  - Bonds in molecules lacking dipole moments will not be detected.
  - The intensity of an absorption band depends on the # of bonds and therefore the strength of bond(s) responsible for the absorption.
  - Bond order affects bond strength, so bond order affects the position of absorption bands.
  - The approximate value can be calculated by Hooke's law.

- The exact position of the absorption band depends on electron delocalization, the electronic effect of neighboring substituents, and hydrogen bonding.
- The predominant effect of the nitrogen of an amide is electron donation by resonance.
- The predominant effect of the oxygen of an ester is inductive electron withdrawal.
- The position and the breadth of the O–H absorption band depend on the concentration of the solution. It is easier to stretch an O–H bond if it is hydrogen bonded.
- The strength of a C–H bond depends on the hybridization of the carbon.
- Interference can be constructive or destructive and leads to overtones and Fermi resonances for ketones, anhydrides, vinyl esters.
- Tells about functional groups by asymmetric bond motion.

1st order stretching frequencies are the easiest to understand by Hooke’s law confirms the presence of aldehydes, ketones, acids and their derivatives, alkenes, alkynes, aromatics, nitro, halides, amines, alcohols for monitoring functional group inter-conversions (experiments 6-9).

- Alcohols and amines display strong broad O–H and N–H stretching bands in the region 3400–3100 cm. The bands are broadened due to hydrogen bonding and a sharp ‘non-bonded’ peak can often be seen at around 3400 cm. N–H’s are usually sharper than O–H’s because of less H-bonding. Acyclic 2° amides may have two stretches.

<table>
<thead>
<tr>
<th>Bond</th>
<th>Wavenumber (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B–H</td>
<td>240</td>
</tr>
<tr>
<td>C–H</td>
<td>300</td>
</tr>
<tr>
<td>N–H</td>
<td>340</td>
</tr>
<tr>
<td>O–H</td>
<td>360</td>
</tr>
<tr>
<td>F–H</td>
<td>400</td>
</tr>
<tr>
<td>Al–H</td>
<td>1750</td>
</tr>
<tr>
<td>Si–H</td>
<td>215</td>
</tr>
<tr>
<td>P–H</td>
<td>235</td>
</tr>
<tr>
<td>S–H</td>
<td>257</td>
</tr>
<tr>
<td>Cl–H</td>
<td>289</td>
</tr>
<tr>
<td>Ge–H</td>
<td>230</td>
</tr>
<tr>
<td>As–H</td>
<td>265</td>
</tr>
<tr>
<td>Se–H</td>
<td></td>
</tr>
<tr>
<td>Br–H</td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>230</td>
</tr>
<tr>
<td>H</td>
<td>265</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type of bond</th>
<th>Monosaccharide</th>
<th>Monosaccharide</th>
<th>Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₂</td>
<td>3200–3210</td>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>CH₃</td>
<td>2900–2910</td>
<td>medium to weak</td>
<td></td>
</tr>
<tr>
<td>C=O</td>
<td>1750–1760</td>
<td>strong</td>
<td></td>
</tr>
<tr>
<td>C–C</td>
<td>1450–1490</td>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>C–N</td>
<td>1500–1650</td>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>C–O</td>
<td>1450–1490</td>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>C–H</td>
<td>3000–3300</td>
<td>very broad</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Compound</th>
<th>Bond</th>
<th>Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetic acid</td>
<td>CH₃</td>
<td>weak</td>
</tr>
<tr>
<td>Methylamine</td>
<td>NH₂</td>
<td>strong</td>
</tr>
<tr>
<td>Methanol</td>
<td>OH</td>
<td>medium</td>
</tr>
<tr>
<td>Water</td>
<td>OH</td>
<td>strong</td>
</tr>
</tbody>
</table>
• Alkene and alkyne C-H bonds display sharp stretching absorptions in the region 3100-3000 cm⁻¹. The bands are of medium intensity and are often obscured by other absorbances in the region (i.e., OH).

• Triple bond stretching absorptions occur in the region 2400-2200 cm⁻¹. Absorptions from nitriles are generally of medium intensity and are clearly defined. Alkynes absorb weakly in this region unless they are highly asymmetric; symmetrical alkynes do not show absorption bands.

• Carbonyl stretching bands occur in the region 1800-1700 cm⁻¹. The bands are generally very strong and broad. Carbonyl compounds which are more reactive in nucleophilic addition reactions (acyl halides, esters) are generally at higher wave number than simple ketones and aldehydes, and amides are the lowest, absorbing in the region 1700-1650 cm⁻¹. More pπ character leads to higher the frequency.

• Carbon-carbon double bond stretching occurs in the region around 1650-1600 cm⁻¹. The bands are generally sharp and of medium intensity. Aromatic compounds will typically display a series of sharp bands in this region.

• Carbon-oxygen single bonds display stretching bands in the region 1200-1100 cm⁻¹. The bands are generally strong and broad. You should note that many other functional groups have bands in this region which appear similar.
Mass Spec.

- Mass spec does not readily distinguish isomers but is good for distinguishing FGI and limited C—C connectivity and the FW and elemental composition.
- The peak with the highest m/z value represents the molecular ion (M)
- Peaks with smaller m/z values are called fragment ion peaks and represent positively charged fragments of the molecule
- Nominal molecular mass: the molecular mass to the nearest whole number. In calculating the molecular masses of molecular ions and fragments, the atom mass of a single isotope of an atom must be used.
- Each m/z value is the nominal molecular mass of the fragment.
- The base peak is the peak with the greatest intensity due to its having the greatest abundance. Usually signifies the most facile cleavage breaking weak bonds break in preference to strong bonds
- Peaks that are attributable to isotopes can help identify the compound responsible for a mass spectrum

Common Cleavages

- Alpha cleavage
- Beta cleavage
- Abstraction
- Cyclization
Simple alkanes tend to undergo fragmentation by the initial loss of a methyl group to form a \((M-15^+)\) species. The carbocation then undergoes stepwise cleavage down the alkyl chain, expelling neutral two-carbon units (ethylene). Branched hydrocarbons form more stable secondary and tertiary carbocations, and these peaks will tend to dominate the mass spectrum.

\[
\begin{align*}
\text{[CH}_3\text{CH}_2\text{CH}_2\text{CH}_3]^{+} & \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2^{+} + \text{CH}_3 \\
m/z = 72 & \\
\rightarrow \text{CH}_3\text{CH}_2\text{CH}_2^{+} + \text{H}_2\text{CH}_2 & \text{m/z = 43} \\
\rightarrow \text{CH}_3\text{CH}_2\text{CH}_3^{+} + \text{H}_2 & \text{m/z = 31} \\
\rightarrow \text{CH}_3\text{CH}_2^{+} + \text{H}_3 & \text{m/z = 29} \\
\rightarrow \text{CH}_3^{+} + \text{H}_4 & \text{m/z = 15}
\end{align*}
\]

The fragmentation of the aromatic nucleus is somewhat complex, generating a series of peaks having \(m/e = 77, 65, 63, \) etc. While these peaks are difficult to describe in simple terms, they do form a pattern (the “aromatic cluster”) that becomes recognizable with experience. If the molecule contains a benzyl unit, the major cleavage will be to generate the benzyl carbocation, which rearranges to form the tropylion ion. Expulsion of acetylene (ethylene) from this generates a characteristic \(m/e = 65\) peak.

The predominate cleavage in aldehydes and ketones is loss of one of the side-chains to generate the substituted oxonium ion. This is an extremely favorable cleavage and this ion often represents the base peak in the spectrum. The methyl derivative (CH\(_3\)CO\(^+\)) is commonly referred to as the “acylium ion”.

Another common fragmentation observed in carbonyl compounds (and in nitriles, etc.) involves the expulsion of neutral ethene via a process known as the McLafferty rearrangement.

The major cleavage observed for these esters, acids and amides involves loss of the “X” group, as shown below, to form the substituted oxonium ion. For carboxylic acids and unsubstituted amides, characteristic peaks at \(m/e = 45\) are also often observed.

In addition to losing a proton and hydroxy radical, alcohols tend to lose one of the alkyl groups (or hydrogens) to form the oxonium ions shown below. For primary alcohols, this generates a peak at \(m/e = 31\); secondary alcohols generate peaks with \(m/e = 45, 59, 73, \) etc., according to substitution.
Following the trend of alcohols, ethers will fragment, often by loss of an alkyl radical, to form a substituted oxonium ion, as shown below.

\[
\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3 \xrightarrow{e^-} \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3^+ \quad m/z = 57
\]

Organic halides fragment with simple expulsion of the halogen, as shown below. The molecular ions of chlorine and bromine-containing compounds will show multiple peaks due to the fact that each of these exists as two isotopes in relatively high abundance. Thus for chlorine, the $^{35}\text{Cl}/^{37}\text{Cl}$ ratio is roughly 3:1 and for bromine, the $^{79}\text{Br}/^{81}\text{Br}$ ratio is 1:1. The molecular ion of a chlorine-containing compound will have two peaks, separated by two mass units, in the ratio 3:1, and a bromine-containing compound will have two peaks, again separated by two mass units, having approximately equal intensities.

\[
\text{CH}_3\text{CH}_2\text{CH}_2\text{Br} \xrightarrow{e^-} \text{CH}_3\text{CH}_2\text{CH}_2\text{Br}^+ + \text{Br}^- \quad m/z = 122 \\
\text{CH}_3\text{CH}_2\text{CH}_2\text{Br} \xrightarrow{e^-} \text{CH}_3\text{CH}_2\text{CH}_2\text{Br}^+ + \text{Br}^- \quad m/z = 124 \\
\text{CH}_3\text{CH}_2\text{CH}_2\text{Br} \xrightarrow{e^-} \text{CH}_3\text{CH}_2\text{CH}_2\text{Br}^+ + \text{Br}^- \quad m/z = 43
\]

**NMR**

- NMR is good for distinguishing C–C connectivity and total assignment.
- Then chemical shift tells about the immediate environment of the proton (electronegativity and anisotropy).
- The intensity of the signal is proportional to the number of protons; the area of a given peak (the integration) is directly proportional to the number of the responsible proton in the molecule. Integrations are given as simplest whole-number ratios.
- The Coupling tells about the number of adjacent protons and their angle relative to the observed proton, but subject to magnetic equivalence.
- The greater the electron density, the greater this 'shielding' will be, hence nuclei which are in electronic rich environments will undergo transition at a higher applied field (upfield) than nuclei in electron poor environments (downfield).
- The chemical shift of the hydroxyl hydrogen of an alcohol moves further downfield with increasing concentration, hydrogen bonding and acidity.
- Because of their favored hydrogen-bonded dimeric association, the hydroxyl proton of carboxylic acids displays a resonance signal significantly down-field of other functions.
- The rapid OH exchange with the deuterium of heavy water can be used to assign hydroxyl proton resonance signals.
- Coupling constants are independent of the external magnetic field, and reflect the unique spin interaction characteristics of coupled sets of nuclei in a specific structure.

Experimentally, for the $^1\text{H}$ and $^{13}\text{C}$ NMR scale is anchored at zero by the NMR absorptions of the molecule tetramethyl silane ((CH$_3$)$_4$Si) for which the carbons and protons are more highly shielded than most common organic molecules.

Pi-electrons are more polarizable than are sigma-bond electrons and a magnetic field induced pi-electron movements that perturb nearby nuclei. The pi-electrons associated with a benzene ring provide a striking example of this phenomenon. The electron cloud above and below the plane of the ring circulates in reaction to the external field so as to generate an opposing field at the center of the ring and a supporting field at the edge of the ring. This kind of spatial variation is called anisotropy, and it is common to non-spherical distributions of electrons, as well. Regions in which the induced field supports or adds to the external field are said to be deshielded, because a slightly weaker external field will bring about resonance for nuclei in such areas. However, regions in which the induced field opposes the external field are termed shielded because an increase in the applied field is needed for resonance. Shielded regions are designated by a plus sign, and de-shielded regions by a negative sign.
**Experiment 1: Fisher esterification**

**Theory & Background:** Esters are widely found in nature and industry. In this experiment, lauric acid (dodecanoic acid) is converted to ethyl laurate. Lauric acid is representative of a class of molecules called fatty acids. These are long, straight-chain carboxylic acids (C₁₂-C₂₀) found as ester derivatives in oils, fats, and waxes. For example, a component of carnauba wax is CH₃(CH₂)₂₃CO₂(CH₂)₃CH₃. Carnauba wax is found in finer automobile waxes and is exuded by the leaves of the Brazilian wax palm tree. Animal fats are fatty acid esters of 1,2,3-propane-triol, also known as glycerol, and are often referred to as triglycerides. In this experiment, acetyl chloride and ethanol are combined to generate HCl acid esters of 1,2,3-propane-triol, also known as glycerol, and are often referred to as tri-waxes and is exuded by the leaves of the Brazilian wax palm tree. Animal fats are fatty acids. These are long, straight-chain carboxylic acids.

**Theory & Background:**

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**Experiment 1: Fisher esterification**

**Reagents**

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</table>

**Procedure:** To a tared 3.0 mL vial equipped with a spin vane, 70-90 mg of lauric acid was added. To the vial was then sequentially added 1.0 mL of EtOH and 3-4 drops of AcCl. A water condenser affixed with a CaCl₂ drying tube was attached to the 0.35 molar solution of lauric acid. The clear suspension was heated on a hot plate at 120°C for 1 hour. The solution was then permitted to cool to room temperature. The room smelled of laurel bushes and laurel leaf oil.

**Work-Up:** The spin vane was replaced with a boiling stone and the contents of the vial were concentrated to 0.3 mL by heating without a condenser to remove unused ethanol. After cooling to room temperature again, 0.5 mL of diethyl ether was added along with 0.5 mL of 1M NaHCO₃ to give several questions on NMR, IR and MS.

**Refractive Index (N).**

Defined as the relative speed at which light moves through a material with respect to its speed in a vacuum. The index of refraction, N, of transparent materials is defined through the equation shown above. c = 3X10⁸ m/s, which is the speed of light in a vacuum and V is the speed of light in some other medium. Since the speed of light is reduced when it propagates through transparent gasses, liquids and solids, the refractive index of these substances is always greater than 1.0. If the refractive index is 0.0010 below or above the literature value, it indicates that impurities are present.

**The thin film technique:** Obtain a salt plate. Salt plates are stored in desiccators. Ideally, the plate was put away clean, although not all students are considerate enough to clean plates after use. If necessary, clean the plates with a small amount of methylene chloride. Ideally, the plates should be transparent, but quite foggy plates usually give acceptable spectra. Take a background spectra of the clean dry salt plate (4 scans). If your sample is a liquid, use a pipet to place a drop of your DRY unknown liquid on the center of salt plate. If your sample is a solid, use a spatula to place a few DRY crystals on the center of the plate. Add one drop of methylene chloride. Allow to dry, and then take your spectra, which will subtract the background. Print your spectra. Remove the plate from the IR, clean it with methylene chloride and place it in the desiccator or pass it on to the next waiting student. You can interpret directly onto the spectra, if you so choose and affix it to your notebook.

**Quiz ideas**

Give several questions on NMR, IR and MS.
neutralize any remaining acid. After agitation, the aqueous layer (bottom) was removed and the organic layer (top) again washed (2 X 0.5 mL of 1M NaHCO₃) to neutralize any remaining acid. After finally washing with water and brine (aq. NaCl gives better separation than H₂O), the organic layer was evaporated to 0.3 mL (3 X expected volume).

**Purification:** A pipet plugged with a small wad of cotton is filled to 50% volume (½ full) with a slurry of 60 micron silica/CH₂Cl₂ and topped off with a 25% volume of Na₂SO₄ (now ¾ full) and placed over a tared 5.0 mL conical vial. The ethereal solution (0.2-0.3 mL) of supposed ethyl laurate was passed through the column into the tared vial. The column contents were rinsed four times with 0.5 mL of CH₂Cl₂. After evaporation (< 0.1 mL remain) the residue was weighed and the % yield calculated.

**Spectroscopy:** An IR spectrum was obtained of the concentrated residue using the thin film technique. (Compare your experimental data (IR and refractive index) to that of actual data given for ethyl laurate. Obtain and interpret an ¹H-NMR, MS and ¹³C-NMR spectrum from your TA. Include the labeled spectra in your notebook directly after the lab. Submit your entire sample in a tared vial to your TA and be sure that your IR is include in your notebook.)

**Waste Disposal:** Any residual CH₂Cl₂ was put in a halogenated organic waste container. Non-halogenated organic waste (ether, and long chain acid/esters) was put into the non-halogenated waste container: Any silica or Na₂SO₄ loaded pipets were dumped into solid waste and then put into a sharps container. Free solid waste (silica, Na₂SO₄) was put into the solids waste container: All aqueous liquids (acetone, ethanol, water) was disposed of in the sink after neutralization or in the basic or acidic waste containers.

**Mechanism:** Both mechanisms shown below are legitimate arrow pushing for an acid catalyzed esterification. Under acidic conditions, a proton is like a penny in a penny cup, in the sink after neutralization or in the basic or acidic waste containers. Free solid waste (silica, Na₂SO₄) was put into the solids waste container: All aqueous liquids (acetone, ethanol, water) was disposed of in the sink after neutralization or in the basic or acidic waste containers.

**Lecture ideas:** Push spectra concepts, lab techniques and calculations. Mechanisms and reactions are the focus of chem 109abc. Chem 6abc is for teaching spectroscopy and laboratory techniques. However, you can cover the mechanism and the role of the RC(O)Cl. Tell your students to place sand in a hole of the heating block and measure temperature of the block with the thermometer provided in their lab drawer. (TURKEY THERMOMETERS ARE FOR TURKEYS).

**TAs:** Be sure that students have completed their entire table when scoring the pre-lab. Give the MC quiz on spectroscopy while contents are refluxing.

Score the products based on the IR and their (N). Watch-out for duplicate IR’s (automatic zero for offending students). Be sure that students explain the MS and carbon and proton spectra when scoring their notebooks.

Reactions should always be monitored by TLC analysis at three different times, if possible.

**Natural Product Isolation**

If you really wanted ethyl laurate, then you’d steam distill it from laurel leaves. Who knows how to steam distill?

**Real Problems**

Residual ethyl acetate can give a false positive for ester in IR spectra. Evaporate for awhile, use heat if necessary. Watch for acids in the IR (unreacted material)

**Calculating % yield**

First, you must calculate the Theoretical Yield. The theoretical yield is the maximum weight or quantity (in grams) of product that can be expected to be formed from a reaction. This number is also used to calculate the percentage yield (see below). The theoretical yield cannot be calculated until the limiting reagent for a reaction has been determined.

The limiting reagent in a reaction is the reactant added to the reaction vessel in the fewest number of moles, after taking into account the stoichiometry of the reaction equation. To determine the limiting reagent, the first step is to write out the molecular/chemical formula and then calculate the molecular or formula weights for the reactants. The second step is to then calculate the # of moles of each reactant added to the reaction vessel. To calculate the number of millimoles of each reactant, divide the quantity of the reactant (mg) by the molecular or formula weight (mg/mmol). This procedure is made slightly more complicated with a weight percentage. (2% solution = 2 mg of compound A / in 100 mg of solvent B). The weight of the solvent B depends upon its density. The third step is to then calculate the # of moles of each reactant added to the reaction vessel.

The percentage yield is one of the most important calculations to learn in organic chemistry. It is a measure of the efficiency of the reaction procedure, and is determined by dividing the isolated yield by the theoretical yield.
Experiment 2: Sodium hypochlorite oxidation of cyclohexanol

Theory & Background: The oxidation of alcohols to ketones or aldehydes is a common reaction. Have you ever wondered how oxy-clean removes stains? For many years, chromium-based reagents were used. In recent years, however, chromium reagents have been used less and less because of their toxicity. Consequently, a number of alternative oxidants have come into use. Sodium hypochlorite, bleach, is one such example. The weak [\text{O—Cl}] bond, resulting in the formation of the non-toxic chloride ion [\text{Cl}^-], is responsible for this reagent's ability to oxidize. This reaction should be monitored by a three lane TLC.

![TLC Diagram](Image)

Please regenerate this table in your notebook filling in any of the blanks:

| Reagents | Values     | cyclohexanol | corrosive acetic acid | sodium hypochlorite | Product | Work-up
|----------|------------|--------------|----------------------|---------------------|---------|---------
|          | formula    | C₆H₁₂O₂      | C₂H₄O₂               | ClNaO               | C₆H₁₂O  | ag. sodium bisulfite |
|          | equiv      | 1.0          | 1.0                   |                     |         | 1.0 expected 5      |
|          | molecular weight | 100.16 mg/mmol | 60.05 mg/mmol | 74.44 mg/mmol |         |         |
|          | density    | 1041 mg/mL   | 1005 mg/mL           | 1206 mg/mL          | 947 mg/mL | solution |
|          | volume     | 0.0961 mL    | 0.262 mL             | 2.08 mL             | 0.103 mL |         |
|          | mass       | 100.0 mg     | 263.7 mg             | 2508 mg             |         | solution |
|          | mmol.      | 0.9984       | 4.393                | 4.043               | 0.9984 |         |
|          | melting point | liquid at RT | liquid at RT         | liquid at RT        | liquid at RT | solution |
|          | boiling point | 161 °C      | 118 °C               | 111 °C              | 115 °C | solution |

Glassware Set-up:

Procedure: Cyclohexanol (100 mg) and acetic acid (0.264 mL) are combined in 5.0 mL vial equipped with a spin vane. The vessel and its contents are chilled in ice water and a Hickman still head is attached. While stirring, sodium hypochlorite (2.08 mL) is slowly added through the throat of the Hickman still head to the vial using a clean disposable pipet (60 drops) so as not to lose compound on the sides of the still-head. The ice bath is removed and the solution stir for one hour at room temperature. During this time, the sodium hypochlorite and acetic acid undergo reaction to generate the oxidant hypochlorous acid (HOCl).

Work-Up: After stirring for 1 hour, a saturated solution of sodium bisulfite is added in dropwise fashion until the reaction mixture gives a negative KI-starch test. If hypochlorous acid is still present, the KI paper turns blue-black; this should not happen once the reaction is fully quenched. After

Quiz ideas

What is the molarity of a 40 wt% solution of NaOH?
Calculations of a table. (in class), including the mass of sodium hypochlorite!
Where does the water go into a reflux condenser (does not matter, highest point, lowest point).
If we formed 2.0 mmol of product Y (mw: 325) from 750 mg of starting material X (mw: 300), what is our percent yield?
Give three likely MS peaks for ethyl laurate as well as the most significant IR and ¹H-NMR signals.
Purification: A pipet plugged from the top with a small wad of cotton, is filled 1/2 full with alumina and topped with a small amount of anhydrous magnesium sulfate is placed over a 3.0 mL conical vial that has been equipped with a boiling chip and tared (weight empty + chip in this case). The column is packed with ether (2 X 0.5mL) and the solvent eluant is discarded. The cyclohexanone solution is then passed through the column into the tared vial. The column contents are rinsed with ether (0.5 mL) into the same tared vial. After evaporation (< 0.1 mL of residue remains) the vial contents are weighed and the percent yield calculated.

Spectroscopy: An IR spectrum is obtained using the thin-film-technique. (Compare your experimental data (IR) to that of actual data given for cyclohexanone. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectrum from your TA. Include all your analyzed spectra in your notebook. Submit your entire sample in a tared disposable vial to your TA along with its IR for grading.

Waste Disposal: Any residual CH$_2$Cl$_2$ is put in a halogenated organic waste container. Non-halogenated organic waste is placed into the non-halogenated waste container. Any silica or Na$_2$SO$_4$ loaded pipets are dumped into solid waste and then thrown into the sharps container. Loose solid wastes (silica, Na$_2$SO$_4$) are put into the solids waste container. All aqueous liquids (acetone, ethanol, water) are disposed in the sink after neutralization, or placed into the basic or acidic aqueous waste containers.

Mechanism: Both mechanisms use legitimate electron counting. However, it is best to think of oxidations as loss of hydride and reductions as addition of hydride.

Lecture ideas: Keep poudring spectroscopy, but be sure to explain how the a still works and how the column works. Discuss IR [ROH goes to RC(O)R]. Explain how magnesium sulfate works.

TAs: Be sure that students have completed their entire table when scoring the pre-lab. Give quiz during the stirring.

Score the products based on the IR and their (N). Watch-out for duplicate IR’s (automatic zero for the offending students). Be sure that students explain the MS and carbon and proton spectra when scoring their notebooks. Reactions should always be monitored by TLC analysis at three different times, if possible. Plates can be developed with 4:1 Hexanes:EtOAc and stained with I$_2$. The cyclohexanol has 1/2 the R$_0$ of the cyclohexanone.
Experiment 3: Horner-Wadsworth-Emmons addition of benzaldehyde, yielding stilbene

Theory & Background: Olefins are an important class of compounds in organic chemistry. They may be synthesized by several means, of which the Horner-Wadsworth-Emmons reaction is among the most convenient and mild. This is considered to be a modification of the original Wittig reaction, which was responsible for George Wittig receiving the chemistry Nobel prize in 1979. In this reaction, phase transfer catalysis occurs at the interface of the two liquids. The reagents themselves reside in different immiscible solvents. The phosphonate becomes deprotonated and undergoes reaction with benzaldehyde. The reaction should be monitored by a three lane TLC.

Glassware Set-up:

Procedure: To a tared 5.0 mL vial equipped with a spin vane, 70-100 mg of the phase transfer catalyst Aliquat 336 is added. This addition is followed by the sequential addition of benzaldehyde (106 mg) and diethyl benzylphosphonate (251 mg). Hexane (1.5 mL) is added to bring the concentration of the benzaldehyde in solution to 0.67M. A water condenser is affixed to the reaction vessel. While stirring vigorously, aqueous NaOH (1.5 mL, 1M, 40 wt %) solution is added through the top of the condenser. The reaction vessel and contents are heated to reflux (about 100 °C). The progress of the reaction can be monitored by TLC. (Be sure to draw pictures of your developed plates into your notebook)

### Theory & Background:

Olefins are an important class of compounds in organic chemistry. They may be synthesized by several means, of which the Horner-Wadsworth-Emmons reaction is among the most convenient and mild. This is considered to be a modification of the original Wittig reaction, which was responsible for George Wittig receiving the chemistry Nobel prize in 1979. In this reaction, phase transfer catalysis occurs at the interface of the two liquids. The reagents themselves reside in different immiscible solvents. The phosphonate becomes deprotonated and undergoes reaction with benzaldehyde. The reaction should be monitored by a three lane TLC.

### Procedure:

To a tared 5.0 mL vial equipped with a spin vane, 70-100 mg of the phase transfer catalyst Aliquat 336 is added. This addition is followed by the sequential addition of benzaldehyde (106 mg) and diethyl benzylphosphonate (251 mg). Hexane (1.5 mL) is added to bring the concentration of the benzaldehyde in solution to 0.67M. A water condenser is affixed to the reaction vessel. While stirring vigorously, aqueous NaOH (1.5 mL, 1M, 40 wt %) solution is added through the top of the condenser. The reaction vessel and contents are heated to reflux (about 100 °C). The progress of the reaction can be monitored by TLC. (Be sure to draw pictures of your developed plates into your notebook)

### Reagents

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Values</th>
<th>Product</th>
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<tr>
<td>benzaldehyde</td>
<td>C₇H₆O</td>
<td>stilbene</td>
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<td>diethyl benzylphosphonate</td>
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<td></td>
</tr>
<tr>
<td>mmol</td>
<td>(Z) 1011 mg/mL</td>
<td></td>
</tr>
<tr>
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<td>(E) 305 °C</td>
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</tr>
<tr>
<td>boiling point</td>
<td>(E) 305 °C</td>
<td></td>
</tr>
<tr>
<td>mass</td>
<td>(744 mm Hg)</td>
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</tr>
<tr>
<td>mmol</td>
<td>(Z) 82 °C</td>
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<td>(0.4 mm Hg)</td>
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</tr>
<tr>
<td>mass</td>
<td>(N) 1.545</td>
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</tr>
<tr>
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</tr>
<tr>
<td>melting point</td>
<td>(Z) 1.622</td>
<td></td>
</tr>
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</table>

### Techniques

**Phase Transfer Catalysis**

There is a practical aspect of this reaction, aqueous sodium hydroxide, needed to generate the carbanion intermediate, is not miscible with the hexane solvent used to dissolve the phosphonate and benzaldehyde. In the absence of other additives such as carbanion generation would be expected to be very slow. The phase-transfer catalyst Aliquat 336, however, accelerates the reaction greatly, because it is soluble in both organic (long aliphatic side chains) and aqueous (charge) media. In the presence of sodium hydroxide, the chloride ion of Aliquat 336 can be exchanged for hydroxide. This material then diffuses into the hexane layer where it can deprotonate the phosphonate. Eventually the ammonium ion becomes the counterion for diethylphosphate. This species then shuttles back into the aqueous phase where phosphate is exchanged for hydroxide to repeat the process.

Aliquat 336

**Quiz ideas**

Calculations of a table. (in class)

**Where does O–H come in the IR?**

**Where does an unstrained ketone C=O come in the IR?**

**Where does the water go into a reflux condenser (does not matter, highest point, lowest point)?**

**What are is multiplicity of each NMR signal for CH₃CH₂OH?**

**How does Aliquat 336 assist the reaction?**

**Draw the ¹H-NMR spectra for ethyl acetate.**

For experiment 2, what instrumental techniques can be used to follow the progress of the reaction? What would you see?
Work-Up: After 1 hour, the reaction was permitted to cool to room temperature. 0.75mL of CH₂Cl₂ is added to dissolve any solids that precipitate upon cooling. In some instances the CH₂Cl₂ does not collect at the bottom, in which case more H₂O is added. Once salt density has been decreased, the CH₂Cl₂ should reside on the bottom of the container. The aqueous layer (sometimes bottom and sometimes top!!) is removed and the organic layer (sometimes top and sometimes bottom!!) is washed with H₂O (0.5mL X 2). After separation, the organic layer is dried over Na₂SO₄, filtered through a pipet containing a cotton plug (see picture on right bottom) into a tared 25mL Erlenmeyer flask containing a boiling chip, and concentrated with heat. The crude material will typically had a mass between 150mg and 250mg after evaporation.

Purification: The (E)-stilbene is recrystallized from a minimum amount of hot absolute ethanol. Slow evaporation afforded pure crystals. The solid residue is weighed and the % yield calculated.

Spectroscopy: A melting point and an IR are obtained. (Compare your experimental data (IR and melting point) to that of actual data given for (E)-stilbene. Obtain and interpret an ¹H-NMR, MS and ¹³C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Submit your entire sample in a tared vial to your TA along with a copy of your IR.)

Waste Disposal: Any residual CH₂Cl₂ is put in a halogenated organic waste container. Non-halogenated organic waste is placed into the non-halogenated waste container. Any silica or Na₂SO₄ loaded pipets are dumped into solid waste and then thrown into the sharps container. Loose solid wastes (silica, Na₂SO₄) are put into the solids waste container. All aqueous liquids (acetone, ethanol, water) are disposed in the sink after neutralization, or placed into the basic or acidic aqueous waste containers.

Mechanism:

Lecture ideas:
Track the components in the two phases during the course of the reaction. Explain how phase transfer catalyst works. Keep hammering spectroscopy, techniques and calculations. Discuss conversion of wt% into molarity. Explain how to recrystallize and why its important for purification and the use of a seed crystal.

TAs:
CHECKOUT THEIR NOTEBOOKS. Be sure that students have completed their entire table when scoring the pre-lab. Be sure they have interpretation of IR, MS and NMRs written on the spectra and these are in their notebooks. Give your quiz during the stirring. Reactions should always be monitored by TLC analysis at three different times, if possible.

Score the products based amount and purity (30-50)+(30-50). Check melting points if unsure of product quality when compared with the IR.

Thin Layer Chromatography
is a fast, convenient method that chemist use to analyze a composition of a mixture. Consider a reaction between two UV active substances that leads to two UV active products of different polarity. The products will adhere to polar silica with differing degrees and display different RF values. However, an RF is only meaningful if the solvent composition is reported. Polar solvents will carry compounds further than non-polar solvents. The least polar compound moves the furthest.

If this hypothetical reaction is incomplete, then the starting materials will also be evident on the plate. Therefore, every TLC analysis should have a minimum of three lanes: one starting material, a mixture of the starting material, the reaction mixture (T₁/₂), and the reaction mixture.

Pipet Filtration
The trick to a good filtration through a drying agent is a good brine work-up and separation. If the organic layer is wet the pipet will plug as the drying agent becomes its corresponding hydrate.
Experiment 4: Hydroboration-oxidation of an alkene

Theory & Background: The hydroboration-oxidation of alkenes to alcohols has long fascinated organic chemists because of its synthetic usefulness and interesting mechanism. H. C. Brown shared the 1979 Nobel prize for developing this and other boron reactions. The reaction of borane with an unsymmetric alkene can give two regioisomeric products: boron at the most substituted carbon (Markovnikov addition) or at the least substituted carbon (anti-Markovnikov addition). The latter is heavily favored, providing for a regioselective reaction. As long as a B-H bond remains, addition reactions to alkenes continue, eventually affording a trialkylborane. Each addition reaction is a four-centered, four-electron process that is extremely rapid. And, each sequential reaction is slightly more regioselective. A GC will be taken of the product mixture at the conclusion of the experiment to determine the ratio of components.

Glassware Set-up:

Procedure 1: To a dry 5.0 mL conical vial equipped with a spin vane, a Claisen (h) adapter and a charged (CaCl₂) drying tube is added octene (150.8 mg, 0.210 mL) via a clean, plastic syringe equipped with an o-ring. The vial and contents are chilled in ice water (0 °C). While stirring, 1.0 BH₃ in THF (0.496 mL) is carefully added via a clean, non o-ring plastic syringe through the top of the adapter in dropwise fashion (1 drop/sec). Caution is used because this exothermic reaction. After addition is complete, the ice bath is removed and the reaction is stirred for 30 minutes. *Syringes for BH₃ additions may be shared.

Work-Up: The Claisen (h) adapter and drying tube are removed. H₂O (3 drops) is slowly added by pipet to quench any unreacted borane.

Please regenerate this table in your notebook filling in any of the blanks:

| Reagent Values | 1-octene (98%) ignore the 2% | 1M borane in THF a stock solution | H₂O₂ 30% (w/w) active | 3M sodium hydroxide 120g/l | octanol
<table>
<thead>
<tr>
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<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
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<tbody>
<tr>
<td>formula</td>
<td>C₈H₁₈</td>
<td>NA</td>
<td>H₂O₂</td>
<td>NaOH</td>
<td>C₈H₁₈O</td>
</tr>
<tr>
<td>equiv</td>
<td>1.0</td>
<td>0.37 (1.1H-equiv)</td>
<td>2.2</td>
<td>1.0 expected</td>
<td></td>
</tr>
<tr>
<td>MW</td>
<td>112.21 mg / mmol</td>
<td>13.8 mg / mmol</td>
<td>34.14 mg / mmol</td>
<td>40 mg / mmol</td>
<td>130.14 mg/mmol</td>
</tr>
<tr>
<td>density</td>
<td>0.715 g/mL</td>
<td>solution</td>
<td>1.11 g / mL</td>
<td>solution</td>
<td>0.827 g/ mL 827 mg/mL</td>
</tr>
<tr>
<td>volume</td>
<td>0.21 mL</td>
<td>0.3012 mL (10 drops)</td>
<td>0.300 mL (8 drops)</td>
<td>0.211 mL</td>
<td></td>
</tr>
<tr>
<td>mass</td>
<td>150.8 mg</td>
<td>solution</td>
<td>335.3 mg (100.6 mg as H₂O₂)</td>
<td>(combined both products)</td>
<td>(what is the bp of H₂O)?</td>
</tr>
<tr>
<td>mmol</td>
<td>1.34 mmol</td>
<td>0.496 mmol</td>
<td>0.9 mmol</td>
<td>1.34 mmol</td>
<td></td>
</tr>
<tr>
<td>boiling point</td>
<td>122 °C</td>
<td>(what is the bp of THF?)</td>
<td>196 °C (1-octanol) 174-181 °C (2-)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Quiz ideas

What is the M of an 80 wt% solution of NaOH?
Draw what happen in a biphasic reaction.
Give a list of compounds and ask, which could act as phase transfer catalyst.
Depict anticipated changes in the IR for propene going to 1-propanol.
Predict the MS, IR and ¹H NMR peaks for ethyl acetate and benzaldehyde.
How many available hydride(s) are in BH₃?
What is the molarity of a 20% by weight solution of hydrogen peroxide (mw = 34.14, d = 1.11 g/mL) in water?
What physical property traditionally dictates separation in gas chromatography?
What is the limiting reagent in this reaction (Exp 4)?
What instrumental techniques are well suited to distinguishing E from Z stilbene?
Procedure 2: A solution of NaOH (0.3 mL of 3M) and HOOH (0.30 mL of 30% active) are mixed in a separate 3.0 mL conical vial using disposable pipet. The same pipet is not used for both liquids to avoid cross-contamination. This mixture is then slowly added in drop-wise fashion (1 drop/second) to the reaction vessel. The resulting solution is stirred for 15 minutes at room temperature.

Work-Up: H$_2$O (1 mL) and ether (1.0 mL) are slowly added and the vial contents are stirred for 5 minutes. Stirring is ceased and the aqueous layer (bottom) is transferred to a 3.0 mL vial. The aqueous layer is acidified with IM HCl (0.1 mL) and extracted with ether (2 X 1 mL). The organic layers are sequentially transferred to a 5.0 mL vial containing the original organic material. The combined organic extracts are washed with water (0.75 mL) and brine (0.75 mL) and then dried over sodium sulfate. After pipet filtration (remember the cotton plug and small amount of organic extracts are washed with water), the vial is not used for both liquids to avoid cross-contamination. This mixture is then slowly added in drop-wise fashion (1 drop/sec) to the reaction vessel. The resulting solution is stirred for 15 minutes at room temperature.

Gas Chromatography Theory

Gas Chromatography, as the name implies, replaces the moving liquid phase of both column absorption chromatography and thin layer chromatography with a gaseous moving phase. Analytical scale GC involves the syringe injection of a small volume of liquid or gas (0.1 mL) through a rubber septum into a stream of inert gas such as helium, which flows at a pre-adjusted rate. The injection port is maintained at a temperature high enough to vaporize the liquid which is then swept by the helium into a heated metal column (6’ x 1/8") maintained at an appropriate temperature. The column is packed with various solid supports. The components of the sample mixture pass through the column at different velocities depending upon their relative gas-liquid phase partition coefficients. In general, non-polar substrates separate on the basis of boiling points while polar substrates separate more (but not exclusively) on the basis of polarity. After separation the components are analyzed by a detector such as a thermal conductivity cell. If the “run” has been standardized by adding a known amount of analyzed substance, then the area under a peak would proportional to the quantity of the substance present. However, in experiment 4, the areas only provide the relative ratios of products. Area = h (peak height at highest point) x w1/2h (peak width at 1/2 maximum height).

Purification: none.

Spectroscopy: The sample is analyzed by GC and the percent yield is calculated on the basis of the preceding crude product weight and the ratio of analytes as determined by GC. An IR (print two, one for your notebook and one for your TA) is obtained with a basis of the preceding crude product weight and the ratio of analytes as determined by Spectroscopy:

Waste Disposal: IMPORTANT - Syringes and their needles do NOT go in the trash. Return the needle in its capped form to the TA. Any residual CH$_2$Cl$_2$ is put in a halogenated organic waste container. Non-halogenated organic waste is placed into the non-halogenated waste container. Any silica or Na$_2$SO$_4$ loaded pipets are first dumped into solid waste and then the glass is thrown into the sharps container. Loose solid wastes (silica, Na$_2$SO$_4$) are put into the solids waste container. All aqueous liquids (acetone, ethanol, water) are disposed in the sink after neutralization, or placed into the basic or acidic aqueous waste containers.

Lecture Suggestions:

Stress anhydrous conditions! Cover needle safety. Explain GC chromatography, and show how the data is to be reported. For example, the product was analyzed by gas chromatography [4ft x ½ in column, 20% DC-200 Chromosorb, 80-100 mesh, 100 °C, helium flow equal to 60mL/mi]. The observed retention times for 1-octene, 1-octanol, and 2-octanol were found to be x, y, and z minutes respectively. Give the crude yield of material the percent yield of each are computed to be X, Y, Z. Discuss the 30% HOOH.

TA Notes

The borane solution is moisture sensitive, the glassware should be oven dried for 30 minutes. Put the intended vial before beginning your lecture or quiz (you will be using a plastic syringes). Monitor the needles given out and those returned. Students can share their borane syringe. H$_2$O$_2$ bleaches skin and clothes, it turns the KI paper black. Use a disposable pipets or tips to measure it.

The b.p. are 122 °C for 1-octene, 174 °C for 3-octanol, 181 °C for 2-octanol and 196 °C for 1-octanol. The order of the peaks on the GC should be the same on the chromatogram.
Experiment 5: Phenyl Grignard addition to benzophenone

Theory & Background: Grignard reagents (RMgX), named for Victor Grignard, who won the Nobel prize for this discovery in 1921, are very useful reagents because of the nucleophilic character of the carbon bonded to magnesium atom. Victor’s breakthrough came with two discoveries—that an ether solvent was vital and that the reaction must be carried out under stringent anhydrous conditions. With such a nucleophilic carbon species, reactions occur with carbon electrophiles, such as carbonyl compounds, to form C-C bonds. In this experiment, triphenylmethanol is obtained from the reaction of phenylmagnesium bromide and benzophenone. Reaction progress should be monitored with three lane TLC analysis.

Grignard reagents (RMgX)

\[ \text{Br} \quad \text{Mg}^+ \quad \text{Et}_2\text{O} \]

Please regenerate this table in your notebook filling in ALL of the blanks

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<thead>
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<th>Reagents</th>
<th>Values</th>
</tr>
</thead>
<tbody>
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<tr>
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<td></td>
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<tr>
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<tr>
<td>volume</td>
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<td>mass</td>
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<tr>
<td>mmol</td>
<td>0.71 mmol</td>
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<tr>
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<td>-31 °C</td>
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<tr>
<td>boiling point</td>
<td>155-156 °C</td>
</tr>
</tbody>
</table>

Glassware Set-up:

Procedure: A drying tube is built from the body of a syringe by removing the o-ring plunger affixing a needle and packing with a cotton plug and dry CaCl₂. Next, a dry 5.0 mL conical vial [A] is charged with anhydrous ether (3 mL) using the 1mL syringe. The same syringe-needle ensemble can be used throughout this experiment. The oven dry, 3.0 mL conical vial [B], equipped with a spin vane, a Claisen (h) adapter, a septum, and fitted with a filled (CaCl₂) drying tube is charged with freshly scratched or ground magnesium (18 mg). One crystal of iodine and while venting the septum with the syringe drying tube some Et₂O (0.5 mL) is added using the same plastic syringe. This make shift drying tube allows solvent to be added to the previously closed system. Another dry, 3.0 mL tared conical vial [C] is charged with bromobenzene (113 mg, 0.075 mL, 7 drops) and Et₂O (0.5mL) and sealed with a septum cap. The etherial solution of bromobenzene is drawn using the same syringe and make shift drying tube. Reaction progress should be monitored with three lane TLC analysis.
Mechanism:

While the reaction proceeds benzophenone (105 mg, 0.30 mL) is added to the now empty 3.0 mL vial [C] along with anhydrous ether (0.30 mL) from the 5.0 mL vial [A]. The reaction mixture in vial [B] is cooled to room temperature. While stirring, the benzophenone solution prepared in vial [C] is cautiously added over one-minute and may result in a gentle reflux (DO NOT REHEAT). After addition is complete, vial [C] is rinsed with the ether (0.3 mL) from vial [A]. This solution was also added to the reaction vessel vial [B]. The above detailed description should assist in preventing the exposure of reagents or solvent to moisture from humid air.

Work-Up: After stirring for 5 minutes, the Claisen head and drying tube are removed. A large amount of solid is visible. The volume is adjusted to 1.5 mL with the remaining ether from vial [A] and additional if needed. While stirring, 3N HCl (1-1.5 mL) is cautiously added until the solids dissolve. This 3.0 mL vial [B] is capped, shaken, and periodically vented. After phase separation, the organic layer (top) is transferred to the empty vial [A] by pipet. The aqueous phase is extracted with ether (2 X 0.5mL) and added to the organic material in vial [A]. The organic material is washed with brine and saturated sodium bisulfite to remove oxidants such as iodine. The solution is dried over anhydrous sodium sulfate and then filtered through a cotton plugged pipet into a 25 mL Erlenmeyer. A boiling stick is added and the solvent evaporated with heat.

Purification: High boiling, 60–80°C, pet ether (10 mL) is added until the solid dissolved and the solution is transferred to a beaker. The solution slowly evaporates in a bench drawer (2-4 days) to give uniform crystals.

Spectroscopy: The crystals are collected and weighed and the percent yield is calculated. A melting point and a IR are obtained. (Compare your experimental data (IR and melt point) to that of actual data given for triphenylmethanol. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectra from your TA. Include the analyzed spectra in your notebook. Submit your entire sample in a tared vial to your TA for scoring.)

Waste Disposal: IMPORTANT - Syringes and their needles do NOT go in the trash. Return the needle in its capped form to the TA. Any residual CH₂Cl₂ is put in a halogenated organic waste container. Non-halogenated organic waste is placed into the non-halogenated waste container. Any silica or Na₂SO₄ loaded pipets are first dumped into solid waste and then the glass is thrown into the sharps container. Loose solid wastes (silica, Na₂SO₄) are put into the solids waste container. All aqueous liquids (acetone, ethanol, water) are disposed in the sink after neutralization, or placed into the basic or acidic aqueous waste containers.

Lecture Suggestions: Explain the set-up, the purpose of the three vials, their septa, and the homemade drying tube. Cover the purpose of scratching the magnesium and the use of iodine as reaction initiators. Explain that biphenyl is formed in small quantities by SET homo-coupling and how its is removed in two stages. Explain the incompatibility of Grignard reagents with a substrates containing acidic hydrogens. Show the reactions that occur when a Grignard reagent is added to an organic acid, or an alcohol.

TAs: Issue three syringes and needles (deduct tech points and return points when syringes are returned) Give your quiz during the stirring. Score products as usual (60-100). There should be a maximum of two duplicate scores. Reactions should always be monitored by TLC analysis at three different times, if possible.

Grignard Reaction Initiators

Why is the Grignard reaction tricky to initiate? The initial step requires electron transfer from the Mg surface to the alkyl halide leading to the formation of R⁺. This step is limited by diffusion control and the available surface area of active magnesium. However, magnesium turnings are generally covered with surface oxides, which preclude its ability to react with unreactive halides in the absence of initiators or mechanic pretreatment. In addition, adsorbed insulating layers and crystal lattice orientation can affect the heterogeneous reaction rates.

In general magnesium turnings are sufficiently prepared for reactive halides after removal of surface oxides and contaminations by hand with pestle and mortar by bending magnesium strips which cause the crystal lattice dislocations. Among the many usual methods of chemical activation, the addition of anhydrous cerium trichloride, iron trichloride, methyl iodide, iodine, and 1,2-dibromoethane are most common.

Here is a good trick. Scratch some magnesium turnings with a needle and immediately submerge the metal in anhydrous ether (Et₂O) and add one crystal of iodine.

TA Notes

Put the vials and stir bars in the oven before your lecture and quiz. [NO PLASTIC ITEMS]. Scratching the metal with the syringe needle exposes fresh surface. The I⁺ in iodine is readily oxidized to I⁻ exposing fresh surface. Don’t use too much iodine and consume all of the magnesium (1 crystal) starting over may be necessary.

The temperature of the drying oven may drop because it is opened so much. Keep it closed as much as possible. Students must not put a wet apparatus (needles or glassware) in the drying oven unless it is rinsed with acetone and air dried.

Use the mL markings on the vials. Bromobenzene (bring your own) can be measured weight into the mL vial or added with a dry pipet. 7 drops is about 113 mg.
Experiment 6: [4+2] Cycloaddition of a masked butadiene

Theory & Background: Otto Paul Hermann Diels and Kurt Alder won the Nobel prize in 1950 for their discovery and development of cyclohexene synthesis using a [4+2] cycloaddition manifold. The resulting six-membered ring can be found in many natural products. The reaction proceeds with control of regiochemistry and stereochemistry and the required conditions are compatible with a large number of functional groups. Thus, the Diels-Alder reaction is frequently exploited. In this experiment, the dienophile, maleic anhydride, is coupled with 1,3-butadiene generated in situ by heat. A small amount of the resulting cyclohexene product is brominated. The reaction should be monitored by three using TLC analysis.

\[
\text{SO}_2(g) + \text{C}_8\text{H}_{10}\text{O}_2 \rightarrow \text{SO}_2\text{Br}_2
\]

Please regenerate this table in your notebook filling in any of the blanks

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<th>Reagents</th>
<th>Values</th>
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</thead>
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<td>maleic anhydride</td>
</tr>
<tr>
<td>formula</td>
<td>C(<em>8)H(</em>{10})O(_2)S</td>
</tr>
<tr>
<td>equiv</td>
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</tr>
<tr>
<td>MW</td>
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<td>density</td>
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</tr>
<tr>
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<tr>
<td>mmol</td>
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<td>65-66 °C</td>
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<tr>
<td>boiling point</td>
<td>solids</td>
</tr>
</tbody>
</table>

Glassware Set-up:

Procedure: To a clean 5mL conical vial, equipped with a spin vane, is added 3-sulfolene (170 mg) and maleic anhydride (90 mg), and followed by the addition of 0.08mL of xylene (5 drops from a disposable Pasteur pipet). A reflux condenser and a drying tube are affixed to the vial and the mixture is heated to 150-160 °C for 15 minutes in a fume hood. Caution is taken because the sulfur dioxide formed during the reaction is very irritating to the eyes and mucous membranes.

Work-up: The reaction vessel is permitted to cool to the touch. The condenser and drying tube are removed and hexanes (0.5 mL) or ligroin (high boiling petroleum ether) is added. Amorphous crystals should form in an oil. Add just enough diethyl ether to solubilize the uncrystallized oil. A reflux condenser and drying tube are reattached and the mixture is refluxed for one minute. The heating is discontinued, the reflux condenser; spin vane, and drying tube are re-

Techniques

Crystallization and recrystallization is a slow method of purification, but it leads to extremely high purity. However, hydrates X·(H\(_2\)O), can pose a problem. In a nutshell, it is easier to pack similar things in tighter spaces than it is to pack dissimilar things. Crystallization begins in a super-saturated solution, which can be reached by lowering the temperature or evaporating a saturated solution.

Heating Reactions

All heating devises work by the same principles. Among the concepts with which organic chemist must be familiar are thermal load and temperature endpoint over-run. While flask size and contents may vary, the heating sources usually does not. Thus, when heating a small item there exists a greater chance that that you will over-run the desired temperature and it will take a long time fore the sample to return to the desired temperature. The trick is to heat SLOWLY!

Quiz ideas

Calculate the liters of SO\(_2\) produced if 1000mg of 3-sulfolene is heated to 160 °C [At STP 22.4 mL / mmol.]

What is the multiplicity, chemical shift and of vinyl protons Ha and Hb in the product. Are they the same? Now consider Maleic anhydride. What is the coupling constant(s)?
moved, and the mixture is permitted to cool. The sides of the vial were scratched below liquid level with a glass stirring rod and crystals began to form. The crystals were quite large and clung to the vial.

**Purification:** The mother liquor is carefully removed with a Pasteur pipet. The solids are given 1-2 minutes to dry and then scraped onto a sheet of filter paper. The capillary action of the paper draws out residual liquid, helping to further dry the crystals. The mass of the product is determined and the percent yield calculated. The presence of an alkene is then verified.

**Spectroscopy:** The melting point and IR spectra were taken for the initial cyclohexene product and compared with the melting point and IR spectra of the known Diels-Alder adduct. (Obtain and interpret an ¹H-NMR, MS and ¹³C-NMR spectra from your TA. Include the labeled spectra in your notebook. Submit your entire sample in a tared vial to your TA for scoring.)

**Waste Disposal:** IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH₂Cl₂ was put in a halogenated organic waste container. Non-halogenated organic waste was put into the non-halogenated waste container. Any silica or Na₂SO₄ loaded pipets were dumped into solid waste and then put into a sharps container. Free solid waste (silica, Na₂SO₄) was put into the solids waste container. All aqueous liquids (acetone, ethanol, water) was disposed of in the sink after neutralization or in the basic or acidic waste containers.

**Mechanism:**

**Lecture Suggestions:**

Explain the importance of s-cis vs. s-trans for the reaction. (What does the s- stand for?) Consider [4+2] transition states, normal vs. inverse demand from the perspective of the components. Cover regiochemistry, if you feel that you can explain it adequately; Most of these topics are ignored in 109abc.

**TAs:** Revisit NMR and discuss the electronegativity and carbonyl anisotropy. Show that ¹H signal for protons alpha to carbonyls and double bonds are in the 2-3.5 ppm range, while vinyls are in the 4.5-6.5 ppm range. (Bisallylic 3.5-5.5 ppm) Revisit IR and carbonyl frequencies for anhydrides. Begin with an anhydrides and work through the carbonyls to cyclohexanones. Then start with cyclohexanone and work to a ketenes. Explain how the bromine solution can be used to quantitate the purity of the 10 mg of crystals (0.0658 mmols). Calculate the molarity of a 2% v/v solution of Br₂ in CCl₄. Reactions should always be monitored by TLC analysis at three different times, if possible.
Experiment 7 Multi-Step (1) Synthesis of diphenylacetylene from stilbene

Theory & Background: Multi-step organic synthesis is extremely challenging. It is the process by which most pharmaceuticals are made. E. J. Corey won the 1990 Nobel prize for his development of the theory and methodology related to multi-step organic synthesis. By combining products from the next three experiments, you will build hexaphenylbenzene. Today is the first part of this process. You will begin by bis-brominating stilbene and then causing a bis-elimination to make diphenylacetylene. Reactions should always be monitored using a three lane TLC for analysis at three different times, if possible.

Please regenerate this table in your notebook filling in any of the blanks

<table>
<thead>
<tr>
<th>Reagent Values</th>
<th>stilbene</th>
<th>pyridinium perbromide</th>
<th>dibromide</th>
<th>85% solid potassium hydroxide</th>
<th>acetylene</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td>C_{14}H_{12}</td>
<td>C_{5}H_{6}Br_{3}N</td>
<td>C_{14}H_{12}Br_{2}</td>
<td>KOH</td>
<td>C_{14}H_{10}</td>
</tr>
<tr>
<td>equiv</td>
<td>1.0</td>
<td>1.0 expected</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>mass</td>
<td>200 mg</td>
<td>400 mg</td>
<td>200 mg</td>
<td>100 mg</td>
<td>—</td>
</tr>
<tr>
<td>mmol.</td>
<td>1.11 mmol</td>
<td>1.25 mmol</td>
<td>1.11 mmol</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>melting point</td>
<td>124 °C</td>
<td>236-237 °C</td>
<td>0.588 mmol</td>
<td>1.49 mmol</td>
<td>0.588 mmol</td>
</tr>
<tr>
<td>boiling point</td>
<td>solids</td>
<td>solids</td>
<td>solids</td>
<td>170 °C 19 mm Hg</td>
<td>—</td>
</tr>
</tbody>
</table>

Glassware Set-up:

**Procedure I:** To a clean 5 mL conical vial, equipped with a spin vane, is added (E)-stilbene (200 mg) and acetic acid (3 mL). A reflux condenser and a drying tube is attached and the mixture is heated to 120-130°C until a homogenous solution forms. The drying tube is removed and, while stirring, pyridinium perbromide (400 mg) is added through the throat of the condenser in one portion. Residual solids are washed into the vial with 1 mL of acetic acid. The material is then heated to reflux (120-130°C) for 5 minutes.

**Work-Up I:** The reaction vessel is permitted to cool to the touch and the condenser and is removed. Crystals form. These are dislodged from the vial with a spatula and then collected by vacuum filtration using a Büchner funnel. The crystal cake is washed with

**Techniques**

**Heating a reaction, cont.**

Heating has solved many a problem for organic chemists. But, heating is not always the best remedy for a sluggis reaction, because a products or reagents can decompose or unwanted side reactions can begin to predomi-nate.

It is important to remember to never heat a closed system unless it is suita-bly protected. A sealed apparatus for this purpose are aptly named “bomb reactors” because of the possibility for a catastrophic explosion.

In general the rate of reaction doubles for every 10 °C rise in temperature.

**Quiz ideas**

What is a chemical test for the presence of an alkene?

Give a GC quiz for computing % yield.

Calculate the molarity of the starting material in acetic acid.

Why was sulfolene used in the preceding experiment rather than buta-diene?

Describe the differences in the MS, IR and C and H NMR the differences between cyclohexenone and cyclohexanol.

What is the function of the KOH in experiment 7?

Why was xylenes used as the solvent in experiment 6?
cold d.i. water (2 mL X 2) and with cold acetone (2 mL X 1). The crystals are set aside to dry.

Purification I: none.

Procedure II: All of the remaining dibromide is transferred to a clean, tared pyrex (13X100mm) test tube. Triethylene glycol is added to make a 0.58M solution. Three equivalents of 85% potassium hydroxide are then added. The mixture is heated for 6-10 minutes using an aluminum block pre-heated to 160-170 °C.

Work-Up II: The test tube is permitted to cool to the touch and 2 mL of d.i. water is added for each 1.0 mL of triethylene glycol. The product precipitates and is collected by vacuum filtration. The crystals are washed with cold 70% ethanol (1 mL X 2) and dried briefly.

Purification II: The product is recrystallized from warmed 95% ethanol (aim to make a 0.5M solution and then supersaturate the solution). If no crystals form, you can add water to the ethanol solution to precipitate, but this fine material is hard to collect by filtration. Large crystals can be collected by vacuum crystallization.

Spectroscopy 1&2: The percent yield is calculated. A melting point and an IR are obtained. (Compare your experimental data (IR and melting point) to that of actual data given for diphenylacetylene. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Do not submit your product to the TA; you will use 150 mg of it in a future experiment. If you failed to recover 150 mg, submit a request in writing to your TA for an exact amount of additional starting material so that you will have 150 mg at the beginning of the next experiment.

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH2Cl2 is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na2SO4 loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na2SO4) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Mechanism:

Lecture Suggestions: Explain why this per-bromide is being used rather than the old stand-by (Br2). Cover the mechanism of bromination. Discuss the importance of adjusting the table of reagents for procedure II based on the yields from procedure I. Cover what 85% KOH really means. Discuss making the 0.58 M glycol solution.

TAs: Please read the note at the right. This product scoring method applies to all of the multi-step experiments (experiments 7-10, 14-15). Please note that students can get a (0) for a product grade in these experiments. Reactions should always be monitored by TLC analysis at three different times, if possible.

Retrosynthetic Analysis

benzaldehyde ↔ hexaphenylbenzene

The arrow means built from. It points in the reverse direction from the conventional synthesis arrow.

Convergent versus linear synthesis

Consider a synthesis that involves 5 steps; each step occurs in 90% yield. The yields at each stage would be as follows (i.e. 90% x 90% at stage 2, etc.). Thus the overall yield of product after 5 steps is 59%.

\[
\begin{array}{ccccccc}
A & B & C & D & E & F \\
90\% & 90\% & 90\% & 90\% & 90\% & 90\% \\
59\% overall
\end{array}
\]

However we could also have run this sequence in a convergent manner and constructed F in a more efficient manner.

\[
\begin{array}{ccccccc}
A & B & C & D & E & F \\
90\% & 90\% & 90\% & 90\% & 90\% & 90\% \\
72\% overall
\end{array}
\]

Compute the overall yield of hexaphenylbenzene. What would the overall yield be if the yield be if stillbene converted to the dibromide in 25%?

\[
\begin{array}{ccccccc}
benzaldehyde & benzoic acid & stilbene & dibromide & furanone & acetylene \\
10\% & 10\% & 10\% & 10\% & 10\% & 10\%
\end{array}
\]

TA Notes

Computing a grade for the product for experiments 7-9. It works sort of like bowling. The product grade depends on how much starting the student request for the experiment where it is used. Use this formulat to compute their score [100% - ((request/150)*100) = score]. Therefore, if a student request 150mg they get a product score of 0%. If they request nothing they get a product score of 100%.

At the end of this lab pass out a sheet of your student’s names so they can indicate what they need for experiment 10. During free time in the next lab you weigh-out their requests for diphenyl acetylene and give it to them, scoring them appropriately.
Experiment 8: Multi-Step (2) Umpolung synthesis of benzoin from benzaldehyde

Theory & Background: Umpolung, or polarity reversal, is a fundamental biological and laboratory transformation. Vitamin B1 causes pyruvate to undergo decarboxylation. It is a surrogate of the polarity reversing reagent, thiamine, that you will use today. Every cell in the body requires vitamin B1 to perform this reaction, which generates adenosine triphosphate (ATP), the fuel the body runs on. In this experiment, you will use thiamine to turn benzaldehyde, an electrophile that undergoes reaction with the nucleophile phenylmagnesium bromide, into a nucleophile, whereupon benzaldehyde will add to itself to make benzoin. The purity of benzaldehyde for this reaction is critical. It must be free of benzoic acid. Your TA has pre-distilled it for you. Reactions should always be monitored by TLC analysis at three different times, if possible.

![Reaction diagram](image)

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Values</th>
<th>benzaldehyde</th>
<th>thiamine + HCl + H₂O</th>
<th>95% EtOH</th>
<th>3M NaOH</th>
<th>benzoin</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td>C₇H₆O</td>
<td>C₂H₅O</td>
<td>NaOH</td>
<td>C₁₄H₁₃O₂</td>
<td></td>
<td></td>
</tr>
<tr>
<td>equiv</td>
<td>1.0</td>
<td>(excess)</td>
<td>40 mg/mmol</td>
<td>212 mg/mmol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MW</td>
<td>106 mg/mmol</td>
<td>337.3 mg/mmol</td>
<td>46 mg/mmol</td>
<td>212 mg/mmol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>den.</td>
<td>1043 mg/mL</td>
<td>solids</td>
<td>789 mg/mL</td>
<td>solids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>vol.</td>
<td>0.4 mL (19 drops)</td>
<td>solids</td>
<td>0.75 mL</td>
<td>0.125 mL</td>
<td>solids</td>
<td></td>
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<tr>
<td>mass</td>
<td>417 mg</td>
<td>65 mg</td>
<td>solution</td>
<td>200 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol.</td>
<td>3.94 mmole</td>
<td>(calcld. as NaOH)</td>
<td>0.94 mmol</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>m.p.</td>
<td>liquid</td>
<td>liquid</td>
<td>solution</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>b.p.</td>
<td>179 °C</td>
<td>solids</td>
<td>solution</td>
<td>solids</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Please regenerate this table in your notebook filling in any of the blanks.

Glassware Set-up: Procedure: To a 5 mL conical vial with spin vane add the powdered thiamine hydrochloride and 0.2 mL of water. Then, add 0.75 mL of 95% ethanol and 0.125 mL of 3M sodium hydroxide and stir. Then add 0.4 mL of freshly distilled benzaldehyde, stir and heat the reaction mixture to 60 °C for 1.5h hours. Do NOT let the temperature go above 65°C at any time during the experiment as this causes undesirable side reactions. The reaction can be analyzed at its supposed conclusion using TLC (3:1 hexanes:EtOAc) by developing the eluted plate in a iodine chamber or with iodine impregnated silica.

Quiz (8) ideas

IR questions on alcohols and ketones would be good. Would diphenyl acetylene show a C=C stretch in the IR? What about trans stilbene? What does the German word Umpolung mean? How does this apply to benzaldehyde? Speculate on the shape of the OH peak of benzoin, if taken in a solution cell? What is its shape? How might the presence of benzoic acid (a) affect the pH of the reaction and (b) affect the crystallization of benzoin at the end?

Techniques

The Seven Steps to Enlightened Distillation
1. Select the drying agent from which the substrate will be distilled.
2. Select the heat source (mantle, Bunsen burner, steam bath, water bath, or aluminum block).
3. Assemble the a clean, dry distillation apparatus from the bottom up. Place heat source. Clamp distillation flask. Set approximate height of receiving flask using a utility clamp. Place condenser into position and secure with joint clamps. Attach tubing to water inlet (lowest) and water outlet (highest) of the condenser. Adjust the height of thermometer (bulb below condenser). Inspect to ensure no joint is under stress, and that the system can be safely heated. (i.e., it is open to the air, nitrogen or vacuum, it is not a BOMB.)
4. Turn on the cold water supply to the condenser. Check for water leaks.
5. Add the drying agent and liquid to be distilled to the pot with boiling stones or a stir-bar.
6. Heat the liquid and collect the product in the receiving flask.
7. Allow the apparatus to cool and disassemble it. Clean all glassware parts thoroughly with acetone (discard in organic wastes) before washing with soapy water.
Work-Up: Upon completion (TLC or 2h), cool the reaction mixture in an ice bath. If crystals do not appear, use a glass rod to scratch the inside surface of the test tube. If this fails to initiate crystallization, then add a few drops of water. The crystals are collected by vacuum filtration with a Büchner funnel. Wash with an ice cold 1:1 mixture of ethanol and water. The washings should remove all of the yellow color and the final product should be colorless. The melting point of the pure product is 134-135°C. If the melting point of your product has a range greater than 4° or deviates much from the 134-135°C range, recrystallize the product from a minimum amount of 95% ethanol (7 mL/g of product). Use the clean dry; crystals for NMR, IR or other spectral analysis, as required.

Purification: none.

Spectroscopy: The percent yield is calculated. A melting point and IR is obtained. (Compare your experimental data (IR and melting point) to that of actual data given for diphenylacetylene. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Do not submit your product to the TA, you will use 200 mg of benzoin in a future experiment. If you failed to recover 200 mg, submit a request to your TA for additional starting material so that you will have 200 mg at the beginning of the that experiment.)

Waste Disposal: Any residual CHCl3 was put in a halogenated organic waste container. Non-halogenated organic waste was put into the non-halogenated waste container. Any silica or Na2SO4 loaded pipets were dumped into solid waste and then put into a sharps container. Free solid waste (silica, Na2SO4) was put into the solids waste container. All aqueous liquids (acetone, ethanol, water) was disposed of in the sink after neutralization failed to initiate crystallization, then add a few drops of water. The crystals are collected by vacuum filtration with a Büchner funnel. Wash with an ice cold 1:1 mixture of ethanol and water. The washings should remove all of the yellow color and the final product should be colorless. The melting point of the pure product is 134-135°C. If the melting point of your product has a range greater than 4° or deviates much from the 134-135°C range, recrystallize the product from a minimum amount of 95% ethanol (7 mL/g of product). Use the clean dry; crystals for NMR, IR or other spectral analysis, as required.

Mechanism:

The second biggest problem is getting crystals. The crystallization procedure used here is referred to as solvent-pair crystallization. Here, benzoin has a low solubility in water; high in ethanol, so as the water content increases for an ethanol/water solution of benzoin, the benzoin (hopefully) begins to precipitate. Often, however, the resulting reaction mixture is an oil, i.e., a super-cooled liquid. Besides scratching the side of the glass of the container with the mixture, there are several other options that you may follow—sometimes a combination of the following are necessary. 1. Scratch the walls of the container with a glass stir rod. Don’t bear down on the glass so much that you break the stir rod—shards of glass aren’t the goal here. 2. Dip the stir rod into the mixture, let it sit dry until you see some small amount of crystalline or powdery solid on the stir rod. Now, place the container in ice water; and continue to scratch the walls of the container. 3. Try reducing the alcohol content of the mixture by letting it evaporate slowly from an uncovered container between lab periods or boiling away with the aid of some boiling chips. 4. Cool and add little more water. This causes more oil to form. Be sure you know which layer is the oil and which is the water. Take the oil, add just enough ethanol to re-dissolve the oil to give a homogeneous mixture, then try scratching, cooling, add a little more water until the solution just becomes a little cloudy, then let it stand uncovered. Chromatography is as always the last resort.

The Umpolung Process
Any process by which the normal alternating donor and acceptor reactivity pattern of a chain, which is due to the presence of O or N heteroatoms, is reversed. Reactivity umpolung is most often achieved by temporary exchange of heteroatoms (N, O) by others, such as P, S and Se. The original meaning of the term has been extended to the reversal of any commonly accepted reactivity pattern. For example, reaction of R–C% CX (X = halide) as a synthon for R–C%C (i.e. electrophilic acetylene) is an umpolung of the normal more common acetylide, R–C%C (i.e. nucleophilic) reactivity.

Vitamin B1 reverses the inherent polarity of an aldehyde.

TA Notes
Notebooks should have a copy of the three TLC plates taken for at least three different times (30min, 60min, 90min) during the reaction. Check the UV and Stain with I2/silica.

Explain the TLC method and how it is used to monitor a reaction and how silica on the plate slows down polar compounds.

Score the product grade as explained earlier.
Experiment 9: Multi-Step (3) Oxidation of benzoin to benzil

Theory & Background: This reaction uses a solution of ammonium nitrate, in the presence of a catalytic amount of copper (II) acetate, to oxidize benzoin to benzil. Reactions should always be monitored by TLC at three different times, if possible.

\[
\text{HO-} \quad \text{cat } \text{Cu(OAc)}_2 \quad \text{OH} \\
\text{NH}_4\text{NO}_3 \quad \text{O} \\
\text{O} \quad \text{O} \\
\text{C}_14\text{H}_{12}\text{O}_2
\]

Please regenerate this table in your notebook filling in any of the blanks:

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Values</th>
<th>benzoin</th>
<th>0.15 M Cu(OAc)(_2) in HOAc/H(_2)O</th>
<th>NH(_4)NO(_3)</th>
<th>benzil</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td></td>
<td>C(<em>{14})H(</em>{12})O(_2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>equiv</td>
<td>1.0</td>
<td></td>
<td>181.5 mg/mmol</td>
<td>80 mg/mmol</td>
<td>1.0 expected</td>
</tr>
<tr>
<td>MW</td>
<td>212 mg/mmol</td>
<td>181.5 mg/mmol</td>
<td>80 mg/mmol</td>
<td>210 mg/mmol</td>
<td></td>
</tr>
<tr>
<td>vol.</td>
<td>solids (52 drps.)</td>
<td>solids</td>
<td>solids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mass</td>
<td>200 mg</td>
<td>solution</td>
<td>110 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol.</td>
<td>0.94 mmol</td>
<td>0.012 mmol</td>
<td>1.375 mmol</td>
<td>0.94 mmol</td>
<td></td>
</tr>
<tr>
<td>m.p.</td>
<td>137 °C</td>
<td>solution</td>
<td>94-95 °C</td>
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<td></td>
</tr>
<tr>
<td>b.p.</td>
<td>solids</td>
<td>solution</td>
<td>solids</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Glassware Set-up:

Procedure: A hot plate and aluminum block is pre-heated to 150 °C. A clean 5mL conical vial, equipped with spin vane, is then charged sequentially with benzoin (200 mg) a 0.15 M copper acetate acetic acid solution (0.8mL), and ammonium nitrate (110 mg). A gas delivery tube and condenser is attached and the vial was placed in the pre-heated block. The end of the gas delivery tube is immersed in a beaker under water. [Caution: when the reaction cools it may cause suction]

Work-Up: Upon completion, as measured by the expected volume of the gas discharge, the gas discharge tube is removed and the reaction mixture is cooled to room temperature. Water (2.0 mL) is added and the mixture is chilled in an ice bath for 10 minutes. The yellow crystals are collected by vacuum filtration with a Büchner funnel and the filter cake is washed with H\(_2\)O (2 X 2 mL). After drawing air through the crystals for several minutes, further drying is accomplished by blotting the solid dry with filter paper. The filtrate is neutralized with 1 M NaHCO\(_3\), until slightly basic and flushed down the drain.

Techniques

De-gassing solvents: The best way to remove water and oxygen from a solvent is to distill it over an appropriate drying agent (such as sodium). This can sometimes be a lengthy (and dangerous) task. For small amounts, it is more efficient to de-gas the solvent by purging with an inert gas such as N\(_2\) or Ar. 1) Place some activated molecular sieves in a hot round-bottomed flask, and purge with nitrogen until cooled to room temperature. Glass readily adsorbs moisture from the air, so it is important to thoroughly oven (or flame) dry all glassware. The sieves will act as a sponge to pick up water dissolved in the solvent. 2) When the flask has cooled, add the solvent and cap with a rubber septum. Secure the septum with copper wire. 3) Purge the solution by injecting a clean needle through the septum and placing it directly in the solvent. Vent the flask with another needle. You should see bubbles. 4) 15–20 minutes should be sufficient.

Another method of degassing a solution is "freeze-pump-thaw." The solution is cooled to -78 °C (dry ice), and then evacuated with vacuum as it is being frozen (using liquid nitrogen, for example), then vacuum is applied for several minutes. The cold bath is removed and the solvent is allowed to slowly warm, once it becomes a liquid, the vacuum is turned off. Repeat this procedure at least two more times.

Quiz ideas

Copper acetate is the oxidizing agent for oxidizing benzoin to benzil, and only 0.012 mmoles are used for 0.94 mmoles benzoin. How can this be possible?

How many nitrates are needed in this reaction?

What structural features make benzil yellow and benzoin white?

How and why does the color change occur when benzoin is oxidized to benzil?

How much gas will be discharged at stp?
Purification: The benzil is slowly recrystallized from hot ethanol (7mL/g), collected, washed with ice cold ethanol, and air dried.

Test for the presence of unoxidized benzoin: Dissolve about 0.5 mg of crude or purified benzil in 0.5 mL of 95% ethanol or methanol, and add one drop of 10% sodium hydroxide. If benzoin is present, the solution soon acquires a purplish color. If no color develops in 2 to 3 minutes, it indicates that the sample is free from benzoin, add a small amount of benzoin, observe the color that develops, and note that contents are capped and shaken vigorously, the color momentarily disappears; when the solution is then let stand, the color reappears.

Spectroscopy: The percent yield is calculated. A melting point and IR are obtained. (Compare your experimental data (IR and melting point) to that of actual data given for benzil. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Do not submit your product to the TA, you will use 100 mg of benzil in a future experiment.)

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH2Cl2 is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na2SO4 loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na2SO4) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Mechanism: Cu^{2+} catalyzes this progress. Oxidations are best viewed as loss of hydride.

Lecture Suggestions:

Quiz ideas (9)

How many mL of CO is produced from 100mg of TPCPD? What is triton B. What does triton B do? Why not use NaOH? How HO^- is produced in the reaction? How much water? If you benzil is mixed with benzoin, what other signals would you see in the IR? Which aromatic carbon in the 13C has the fewest hydrogens connected to it.

TAs:
Experiment 10: Multi-Step (4) Synthesis of hexaphenylbenzene from benzil

Theory & Background: The multi-step synthesis converges in this experiment as a bis-aldol condensation reaction is followed by a Diels-Alder [4+2] cycloaddition with diphenyl acetylene and concomitant loss of carbon monoxide. You will make a KBr pellet of the first intermediate and record it's IR. Reactions should always be monitored by TLC analysis at three different times, if possible.

O
O
O

Please regenerate this table in your notebook filling in any of the blanks

<table>
<thead>
<tr>
<th>Reagents Values</th>
<th>benzil</th>
<th>dibenzyl ketone</th>
<th>triton B 40 wt% in MeOH (1.79M)</th>
<th>HO(CH_2CH_2O)_3H</th>
<th>tetraphenyl cyclopenta-dienone</th>
<th>diphenyl acetylene</th>
<th>hexaphenyl benzene</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<td>equiv</td>
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<td>1.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
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</tr>
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<td>0.48 mmol</td>
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<td>-</td>
<td>-</td>
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<td>(bp of MeOH)</td>
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<td>solids</td>
<td>solids</td>
<td>solids</td>
</tr>
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</table>

Glassware Setup:

Procedure 1: A hot plate and aluminum block is preheated to 150 °C. To a dry 3.0 mL conical vial, equipped with a spin vane, is added benzil (100 mg), dibenzyl ketone (100mg) and triethylene glycol (0.5 mL). An air condenser is attached. A clamp is attached to the air condenser and the mixture is heated until homogenous. The heat source is removed and Triton B (0.093 mL) is immediately added through the condenser. Upon cooling purple crystals formed.

Work-Up 1: While stirring at room temperature, methanol (1.3 mL /100 mg of benzil) is added. The vial is cooled in an ice bath and the purple crystals are collected by vacuum filtration with a Büchner funnel. The crystals are washed with ice cold methanol until the filtrate is purple-pink, not brown. The crystals are collected and air dried. The filtrate is placed

Quiz (10) ideas

What is the molarity of a 10% by weight solution of hydrogen peroxide (mw = 34.14, d = 1.11 g/mL) in water?

Draw and label the ¹H NMR spectrum of 3-methoxy-pentane. Be sure to place peaks at appropriate chemical shifts in reference to one another, and illustrate multiplicity.

Given the following TLC plate, if you had to choose between the two structures below, which do you think would correspond to the sm spot?

In total synthesis what aspect is the most critical in evaluating the efficiency of a route?

What is the function of the benzyltrimethylammonium hydroxide? Why is it better to use in this experiment than aqueous NaOH?

Techniques

High Pressure Reactors

Hydrogenations are often run a high pressure. The apparatus below, a Parr Shaker, is for performing reactions at 10-100psi.

Higher pressures 100-5000 psi are obtained using a Parr bomb.
in an organic waste container. The percent yield of tetra-phenylcyclopentanone is calculated.

**Purification:** If the crystals are not well formed or if the melting point is low, then place the material in a reaction tube, add 0.6 mL of triethylene glycol, stir with a thermometer (You will need to use a mercury thermometer from one of the melting point apparati to do this. Please remember to put it back), and raise the temperature to 220 °C to bring the solid into solution. Let it stand for crystallization (if initially pure material is recrystallized, the recovery is about 90%). Save 10 mg of these crystals for measuring a melting point and taking a KBr pellet IR. Run all of the remainder in the next reaction. If you need a supplement, ask your TA.

**Procedure 2:** To a pyrex tube is added the recovered tetra-phenyl-cyclopentanone (150 mg), diphenylacetylene (150 mg), [if you have more of either; run what you have, supplement from the TA up to 100 mg] and silicone oil (1mL). The actual amounts used are recorded. Care is taken to ensure that the components are completely dry before mixing (else, spattering could occur). The mixture is brought to a boil (250 °C) by rotating the pyrex test-tube over a micro-burner. A colored, homogeneous solution results after heating for 1-3 minutes. Heating is continued for an additional 10 minutes and a tan solid appears.

**Work-Up 2:** After cooling to room temperature, the mixture is diluted with hexanes (3.0mL). The crystals are collected by vacuum filtration with a Büchner funnel. The filter cake was washed with hexanes (2 X 1 mL) and cold toluene (2 X 1 mL). The filtrate is placed in an organic waste container. The cake is air dried for several minutes.

**Purification:** The crystals are further purified by recrystallization from a minimum amount of hot diphenyl ether (259°C).

**Spectroscopy:** The percent yield is calculated. The melting point of last product is too high to measure. KBr pellet IR spectra are obtained for the two products. (Compare your experimental data (IR) to that given for tetraphenylcyclotetradione and hexaphenylbenzene. Obtain and interpret an ¹H-NMR, MS and ¹³C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Submit your entire sample in a tared vial to your TA for scoring.

**Waste Disposal:** IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH₂Cl₂ is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container: Any silica or Na₂SO₄ loaded pipets are dumped into solid waste and the glass placed in a sharps container: Free solid waste (silica, Na₂SO₄) is put into the solids waste container: All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

**Mechanism:**

**Lecture Suggestions:** Cover an aldol condensation and a Diels-Alder reaction. Note however that most 6B/109B students have only covered a simple [4+2] Diels-Alder reaction and have not yet seen an aldol reaction. You should not quiz on the mechanisms. Demonstrate a KBr pellet formation.

**TAs:** The final product is scored as usual.
**Experiment 11: Acetanilide synthesis and aromatic bromination**

**Theory & Background:** Bromination of aniline is an uncontrollable reaction. In this sequence, however, aniline is first acetylated. It thereby becomes less reactive. Bromination then proceeds in a regioselective manner. Reactions should always be monitored by TLC analysis at three different times, if possible.

\[
\text{NH}_2 + \text{HCl} \rightarrow \text{NH}_2\text{Cl} \\
\text{NH}_2\text{Cl} + \text{Ac}_2\text{O} \rightarrow \text{NH}_2\text{C}____ \text{Ac}_2 \\
\text{NH}_2\text{C}____ \text{Ac}_2 + \text{Br}_2 \rightarrow \text{NH}_2\text{C}____ \text{Br}_2 \\
\text{NH}_2\text{C}____ \text{Br}_2 + \text{NaHSO}_3 \rightarrow \text{NH}_2\text{C}____ \text{H} + \text{Na}^+ + \text{SO}_4^{2-}
\]

Please regenerate this table in your notebook filling in any of the blanks.

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<th>Values</th>
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</thead>
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</tr>
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</tr>
<tr>
<td>boiling point</td>
<td>184 °C</td>
</tr>
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**Glassware Set-up:**

**Procedure:** To a 13 x 100-mm test tube is added aniline (0.1mL), and followed by the addition of water (0.5 mL) and 12M hydrochloric acid (0.1 mL, 3 drops). This converts the aniline into its hydrochloride salt. A pellet of charcoal (20 mg Norit) is added. The charcoal removes compounds that result from oxidation of aniline, which gives color to the solution. After swirling for 3-5 minutes, the suspension is filtered through a cotton plugged pipet into a 3.0 mL conical vial containing a magnetic spin vane (may still be yellow). The residue remaining in the test tube is rinsed with water (0.5mL) and filtered into the 3.0 mL conical vial. Sodium acetate-trihydrate (150 mg) is added to the rinsed 13 x 100-mm test tube, followed by the addition of d.i. water (0.5 mL). This sodium acetate solution is set aside. To the 3.0 mL conical vial containing the aniline hydrochloride is added acetic anhydride (0.15mL, 10 drops) and the solution of sodium acetate. The sodium acetate, being basic, frees the aniline’s lone pair of electrons from the salt and permits its subsequent reaction with the acetic anhydride. The reaction is rapid and a precipitate forms. Stirring is continued for (5 min) to ensure mixing of the reagents and reactant. The 3.0-mL conical vial is then placed in an ice-water bath for 10 minutes to induce crystallization of product.

**Techniques**

**Quiz ideas**

How does one evaluate a total synthesis?
Predict the IR of acetanilide? How many signals for the N-H.
Explain significant peaks in the Mass Spec.
How many signals in the 1H-NMR for 4-bromo-acetanilide?
Show what the MS would look like for the molecular ion.
What is the purpose of the NaHSO3 quench?
Work-Up: The product is isolated by vacuum filtration with a Büchner funnel. The 3.0-mL conical vial is rinsed with water (0.5mL x 2). These rinsings are used to wash the crystals. The crystals are blotted onto a paper towel to accelerate air drying. The aqueous filtrate is disposed down the drain after dilution with water. The percent yield of acetanilide is calculated. About 10 mg of this material was saved for spectra and m.p. analysis.

Procedure 2: In a fume hood, a 5.0-mL conical vial, is equipped with a cap and charged with acetanilide (100 mg) and glacial acetic acid (16 drops). Gentle agitation results in dissolution of the acetanilide. Because bromine is a powerful irritant and oxidant and displays a significant vapor pressure and extreme care should be exercised in the next step. Once acetanilide is dissolved, the 7M Bromine/HOAc solution (12 drops) is added to the conical vial, which is then capped. The reaction mixture is gently agitated periodically. After 10 minutes, water (1.5 mL) is added to the reaction mixture, followed by aqueous sodium bisulfite (20 drops) [crystals form]. The red-orange color should dissipate due to the reduction of the bromine [Br+\textsuperscript{+}] by the sodium bisulfite.

Work-Up: The crystals are collected by vacuum filtration using a Büchner funnel, washed with cold water (1mL x 3), and partially dried by drawing air through the crystals. Blotting with paper towels accelerates the drying process. The filtrates are flushed down the drain.

Purification: The crude 4-bromoacetanilide is re-crystallized from 95% EtOH.

Spectroscopy: The percent yield is calculated. A melting point and IR spectrum is obtained for acetanilide and 4-bromoacetanilide. (Compare your experimental data (melting point and IR) to that of actual data given for acetanilide and 4-bromoacetanilide. Obtain and interpret an \textsuperscript{1}H-NMR, MS and \textsuperscript{13}C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Submit your entire sample in a tared disposable vial to your TA.)

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH\textsubscript{2}Cl\textsubscript{2} is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na\textsubscript{2}SO\textsubscript{4} loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na\textsubscript{2}SO\textsubscript{4}) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Mechanism:

Lecture Suggestions: Explain the importance of the peptide bond in nature. Explain why rotation is hindered and rotamers are the reason for the two different methyl singlets. Keep pounding spectroscopy (NMR, IR, MS).
Experiment 12: Synthesis of α- and β- glucose penta-acetate

Theory & Background: Sugars and their anomers play important roles in cell recognition.

Glucose exists in solution as an equilibrium between the α and β anomers of its pyranose hemiacetal and its open form. However, the solid starting material provided by Aldrich exists exclusively as α-anomers. When using sodium acetate and acetic anhydride, the less hindered, equatorial hydroxyl group of the β isomer undergoes reaction much much faster than the axial hydroxyl group of the α isomer. This results in the preferential formation of the β-glucose pentaacetate. What can you deduce about the difference in the rate of acylation for the α and β anomers versus the equilibrium constant between them? In the second experiment, you’ll generate an oxonium ion. Because of the anomeric effect, it undergoes reaction much much faster than the α anomer. You’ll note the different NMR spectra. You will need to find a partner so that one sets a hot plate to 110 °C while the other sets theirs to 95 °C. You will share the hot plates, but you’ll each run both experiments.

The anomers of both D-glucose pentaacetate and L-glucose pentaacetate have been found to display insulino tropic potential.

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Glassware Set-up:

**Procedure 1:** A hot plate is pre-heated to 110-120 °C. To a 5mL conical vial, equipped with a spin vane, is added anhydrous glucose powder (100 mg), anhydrous sodium acetate powder (80 mg), and acetic anhydride (0.75 mL, 45 drops). An air condenser and drying tube are attached to the vial. The stirring mixture is heated at 110-120°C for one hour.

**Work-Up 1:** After 1h, heating is discontinued and the cooled vial is poured onto ice (5 mL) in a small beaker. The excess acetic anhydride reacts with water making acetic acid. Acetic acid is fairly water soluble. An oil separates and solidifies after 30 minutes of stirring with a glass rod. The solids are collected by filtration using a Hirsch funnel and washed with cold d.i. water (2 mL).

**Purification 1:** The solid is recrystallized from a minimum amount of hot 95% ethanol (1-2 times). The percent yield is calculated.

**Procedure 2:** The other hot plate (your partner’s) is preheated to 95-100°C. To a 5mL conical vial equipped with a spin vane is added anhydrous glucose powder (100 mg), zinc chloride (30 mg), and acetic anhydride (0.75 mL, 45 drops). An air condenser and drying tube is attached to the vial. The stirring mixture is heated at 95-100°C for 1 hour.

**Work-Up 2:** After 1h, heating is discontinued and the cooled vial was poured onto ice (5 mL) in a small beaker. The excess acetic anhydride reacted with water, making acetic acid. Both acetic acid and zinc chloride are fairly water soluble. An oil separated and solidified after 30 minutes of stirring with a glass rod. The solids were collected by filtration using a funnel and washed with cold d.i. water (2 mL).

The \(^1\)H NMR spectra to the two acetylated glucoses are similar of course, but key differences exist. The proton at C-1 called H-1 has the greatest chemical shift because it is on a C attached to two O’s. The proton H-1 is at 5.7 ppm in the β isomer and at 6.3 ppm for the α isomer. Typically axial protons are at higher field (smaller chemical shift) than are the equatorial protons. Also notice that in the chair conformational structure of the β isomer that H-2 is anti to the H-1, and this geometry results in a doublet with splitting (coupling) of 9.0 Hz. In contrast the α isomer where the H-1 and H-2 are gauche to one another, has a splitting of only 3.5 Hz in accord with Karplus equation. These compounds were among those used in the early days of NMR to study these effects.

**TA Notes**

If there are not enough drying tubes then build one from a Claisen adapter.
Purification 2: The solid is recrystallized from a minimum amount of hot 95% ethanol (1-2 times). The percent yield is then calculated.

Spectroscopy 1&2: Melting point and IR spectra are obtained for both products. (Compare your experimental data (melting point and IR) to that of actual data given for both products. Obtain and interpret an $^1$H-NMR, MS and $^{13}$C-NMR spectrum from your TA. Note the slight differences between the diastereomers particularly in vicinal coupling. Include the labeled spectra in your notebook. Submit your combined samples in a tared vial to your TA. Note the origins of the amounts.)

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH$_2$Cl$_2$ is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na$_2$SO$_4$ loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na$_2$SO$_4$) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Lecture Suggestions: First, re-examine NMR and discuss diastereotopic signals and coupling. Be sure to explain the Karplus rule ($\cos^{-1} 90^\circ = 0$). Then explain the hemiacetal-aldehyde equilibrium. Next, discuss the anomeric effect. It is easiest to explain it as a means to prevent destabilization from electron-electron repulsion in the product.

Techniques

Quiz ideas

Excluding O–H denote the proton that would be the most shielded.

Other than a pyranose, are there other forms of glucose?

Explain the anomeric effect?

What governs a kinetic distribution of diastereomers?

What governs a thermodynamic distribution of diastereomers?

How do these topics apply to experiment 12?

How might you monitor the reaction by IR?

What is the role of zinc chloride?

What does alpha and beta signify in general?

What does alpha and beta signify for carbohydrates?
Experiment 13: Synthesis of methyl red and orange via a diazo-nium coupling

Theory & Background: Today you’ll again be running two experiments in parallel. In the presence of nitrous acid, formed from sodium nitrite and hydrochloric acid, an aromatic amine is converted to an aryl diazonium ion \( \text{Ar}^+\text{N}_2^- \). Close to 0 °C, the aryl diazonium ion is stable for hours. At higher temperature it loses \( \text{N}_2 \), forming a very reactive aryl cation. In these experiments, the respective diazonium ions undergo C–N bond formation with N,N-dimethylaniline. This forms azo-dyes, compounds that are used as acid-base indicators. Reactions as usual, should always be monitored by TLC analysis at different times, if possible.

Procedure 1: Sulfanilic acid (100mg, 0.58 mmoles) is added to a 5 ml conical vial equipped with a spin-vane. If sulfanilic acid monohydrate is used, 110mg is required. A 0.24 M solution of sodium carbonate (1.2 mL, 31 drops) is added and the reaction vessel is heated until all solids dissolved. The vessel is cooled to room temperature and sodium nitrite (40 mg, 0.58 mmoles) is added. 3M hydrochloric acid (0.5 mL, 0.074 mL 4 drops) is then slowly added. The mixture changes color to orange. While stirring, the vial is heated in a beaker of boiling water. Most of the solids dissolve. Upon cooling to 0 °C, crystals form. These are collected by vacuum filtration. The filter cake is washed with saturated brine.

Work-Up 1: After stirring for 10 minutes, a paste forms and 3 M NaOH (1 mL, 24 drops) is then slowly added. The mixture changes color to orange. While stirring, the vial is heated in a beaker of boiling water. Most of the solids dissolve. Upon cooling to 0 °C, crystals form. These are collected by vacuum filtration. The filter cake is washed with saturated brine.

Please regenerate this table in your notebook filling in any of the blanks

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<th>0.24M Na₂CO₃</th>
<th>sodium nitrite</th>
<th>3M HCl</th>
<th>dimethyl aniline</th>
<th>acetic acid</th>
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<th>methyl orange</th>
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</tbody>
</table>

The Enterobacteriaceae, Gram-negative bacilli, are the most frequently encountered bacteria isolated from clinical specimens. Widely dispersed in nature, the Enterobacteriaceae occupy a number of ecological niches including the intestinal tracts of humans and animals. Before the advent of antibiotics infectious diseases caused by the Enterobacteriaceae were well defined. Endotoxic shock is one of the potentially life-threatening consequences of infection by Gram-negative bacteria including the Enterobacteriaceae. Endotoxin is a lipopolysaccharide found in the outer membranes of Gram-negative bacteria. The lipid-A portion of the lipopolysaccharide molecule is primarily responsible for the bioactive properties of endotoxin. Endotoxin is highly antigenic and its structure is highly conserved in all strains of Gram-negative bacilli. Many bacteria use one of two alternative routes for reoxidizing NADH, which generate large quantities of acetic acid, lactic acid, and formic acid and lesser amounts of succinic acid, propionic acid, butyric acid, and butyl alcohol. In olden days, methy red was used to screen surfaces for the presence of bacteria. Today, tests for detecting Gram-negative sepsis are based on the development of monoclonal antibodies specific for the conserved lipopolysaccharide molecule on the surface of the bacteria.
Mechanism

All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Waste Disposal

- Any silica or NaOH is put into the solid waste container.
- Non-halogenated organic waste is put into the non-halogenated waste container.
- All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Techniques

Methyl orange is an intensely colored compound used in dyeing and printing textiles. It is also known as C.I. Acid Orange 52, C.I. 13025, helianthine B, Orange III, Gold orange, and Tropaeolin D [1]. Chemists use methyl orange as an indicator in the titration of weak bases with strong acids. It changes from red (at pH 3.1) to orange-yellow (at pH 4.4).

TA Notes

- Use a KBr pellet for the IR. These dyes stain the salt plate.
- Do not take melting point.

Quiz (13) ideas

- What might happen if aniline were used instead of dimethyl aniline?
- In UV-Visible Spectroscopy, the presence of what lead to bathochromic shift?
- What are some true concerning the absorption of light in UV-Visible spectroscopy?
- What does Hyperchromic means?
- What does the absorption intensity reflect?
- Explain how deprotonation effects the UV-Vis of these compounds.
- Why would these compounds colors be different at differing pH?

Purification 1: Recrystallization from water and calculation of the percent yield.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Values</th>
<th>anthranilic acid</th>
<th>3M HCl</th>
<th>3M sodium nitrite</th>
<th>3M sodium acetate</th>
<th>dimethyl aniline</th>
<th>3M sodium hydroxide</th>
<th>methyl red</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td></td>
<td>HCl</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>HONa</td>
</tr>
<tr>
<td>equiv</td>
<td>1.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.0 exp.</td>
</tr>
<tr>
<td>MW</td>
<td>137</td>
<td>69</td>
<td>82</td>
<td>121</td>
<td>40</td>
<td>269</td>
<td></td>
<td></td>
</tr>
<tr>
<td>density</td>
<td>solids</td>
<td>soln.</td>
<td>solution</td>
<td>solution</td>
<td>956 mg/ mL</td>
<td>solution</td>
<td>solids</td>
<td></td>
</tr>
<tr>
<td>volume</td>
<td>solids</td>
<td>0.7 mL</td>
<td>0.2 mL</td>
<td>0.3 mL</td>
<td>6 drops</td>
<td>0.1 mL</td>
<td>solids</td>
<td></td>
</tr>
<tr>
<td>mass</td>
<td>80 mg</td>
<td>soln.</td>
<td>41 mg</td>
<td>(calc. as NaOAc)</td>
<td>103 mg</td>
<td>(calc. as NaOH)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol.</td>
<td>0.58</td>
<td>2.1</td>
<td>0.6</td>
<td>1.0</td>
<td>0.85</td>
<td>0.3</td>
<td>0.58</td>
<td></td>
</tr>
<tr>
<td>mp or bp</td>
<td>144 °C</td>
<td>soln.</td>
<td>solution</td>
<td>solution</td>
<td>193 °C</td>
<td>solution</td>
<td>181 °C</td>
<td></td>
</tr>
</tbody>
</table>

Procedure 2: Anthranilic acid (80 mg, 0.58 mmole) is added to a 3 mL conical vial equipped with a spin vane. 3M hydrochloric acid (0.7 mL, 17 drops from a Pasteur pipet) is then added. Heat is applied if necessary to dissolve solids. The vial is then cooled (0-5°C) in a small beaker of ice water. In another vial or test tube is added 44 mg (0.64 mmoles) of sodium nitrite, which is then dissolved in water (0.20 mL, 5 drops from a Pasteur pipet). The sodium nitrite solution is cooled in ice water and then the chilled solution is added to the solution of anthranilic acid via a pipet. Dimethylaniline (103 mg, 0.85 mmole, 6 drops from a Pasteur pipet) is then added and the contents are stirred for 15 minutes. Next, freshly prepared 3M NaOAc (0.3 mL) is added and stirring is continued for 20 minutes. The reaction is then warmed to room temperature and then 3 M NaOH (0.1 mL, 3 drops from a Pasteur pipet) is added and the contents are left standing for 30 minutes to ensure thorough precipitation.

Work-Up 2: Methyl red is collected by filtration using a Hirsch funnel. The reaction vial is rinsed with water. The filter cake is washed with 3M acetic acid (about 0.5 mL) to remove excess dimethylaniline. The filter cake is then washed with a small amount of water.

Purification 2: In a fume hood, the product is recrystallized from methanol. The solution is cooled to 0 °C before filtering.

Spectroscopy 1&2: DO NOT TAKE MELTING POINTS. The percent yield is calculated. IR spectra are obtained for both products. (Compare your experimental data (melting point and IR) to that of actual data given for methyl red and orange. Obtain and interpret an 1H-NMR, MS and 13C-NMR spectrum from your TA. Note the differences in couplings for o-substitution verses p-substitution. Include the labeled spectra in your notebook. Submit both of your samples in separate tared vials to your TA.)

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH₂Cl₂ is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na₂SO₄ loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na₂SO₄) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Mechanism: Students should be able to propose one on their own.
Experiment 14: Synthesis of 2-(4-methylbenzoyl)benzoic acid

Theory & Background: Friedel-Crafts acylation involves the addition of a benzene ring at the carbonyl of an acyl chloride or an anhydride. This reaction has several advantages over Friedel-Crafts alkylation. Among these is the fact that the product is less reactive than the starting material, so multiple acylations do not occur. Also, there is little possibility for carbocation rearrangements in the side chain being added. Reactions, as usual, should be monitored by TLC analysis at different times, if possible.

![Chemical structure of 2-(4-methylbenzoyl)benzoic acid]

Please regenerate this table in your notebook filling in any of the blanks:

<table>
<thead>
<tr>
<th>Reagents</th>
<th>phthalic anhydride</th>
<th>toluene</th>
<th>aluminum trichloride</th>
<th>solid water (ice)</th>
<th>2-(4-tolueyl) benzoic acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>equiv</td>
<td>1</td>
<td>7</td>
<td>2.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MW</td>
<td>148 mg/mmol</td>
<td>92 mg/mmol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>density</td>
<td>solids</td>
<td>solids</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>volume</td>
<td>solids</td>
<td>0.75 mL</td>
<td>45 drops</td>
<td>solids</td>
<td></td>
</tr>
<tr>
<td>mass</td>
<td>150 mg</td>
<td>300 mg</td>
<td>300 mg + 300 mg + 300 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol</td>
<td>1.01</td>
<td>7</td>
<td>2.3</td>
<td></td>
<td>1.01</td>
</tr>
<tr>
<td>melting point</td>
<td>131 °C</td>
<td>liquid</td>
<td></td>
<td></td>
<td>138-139 °C</td>
</tr>
<tr>
<td>boiling point</td>
<td>solids</td>
<td>110.6 °C</td>
<td>solids</td>
<td></td>
<td>solids</td>
</tr>
</tbody>
</table>

Glassware Set-up:

Procedure: Phthalic anhydride (150 mg) is weighed into a dry 3.0mL vial, a spin vane is added, and the apparatus is affixed to an air condenser. Toluene (0.75 ml, 45 drops) is then added. The reaction vial is chilled with ice water. Three-full spatulas of fresh AlCl₃ (300 mg) are provided by the TA, directly into the reaction vessel from a freshly prepared small 10g vial. A gas bubbling apparatus, with toluene in the flask, is assembled as shown. The reaction vial is cautiously warmed with hand heat. Reaction speed was gauged by the amount of gas bubbles. If the reaction becomes too vigorous, the vial is cooled with ice water. Warming and cooling is alternated on the hot plate, without heat, until the reaction slows. The reaction is then heated on an aluminum block at the lowest heat setting until gas evolution stops.

Techniques

Quiz (14) ideas

If we have 100mg of glucose (mw = 180.16) in 0.75mL of acetic anhydride (mw = 102, d = 1.08 g/mL), what is the molarity and weight percent of the solution?

What role does AlCl₃ play in this reaction? Explain the C=O stretches in the IR for phthalic anhydride?

What is a Fermi resonance? Speculate on the shape of this RCO₂-H peak in the IR using solution cells?

Where do you think this signal will come in the ¹H-NMR?

Calculate the number of moles of hydrogen chloride liberated in the microscale synthesis of 2-benzoylbenzoic acid. If this gaseous acid were dissolved in water, hydrochloric acid would be formed. How many milliliters of concentrated hydrochloric acid would be formed in this reaction? The concentrated acid is 12 M in HCl.

Write a mechanism for the formation of 2-(4-methylbenzoyl)benzoic acid from toluene and phthalic anhydride using an aluminum chloride catalyst.
Work-up: The gas bubbler is removed and a small piece of ice is added through the condenser to the vial. After reaction subsides, another small piece of ice is added. This process is continued until 1.0g of ice has been added. 3M HCl (1.3 mL, 31 drops) is then slowly added. After cooling to room temperature, ether (b.p. 35°C, 1.5 mL) is added. A screw cap septum is attached and the mixture is well shaken, venting when needed. The aqueous layer (bottom layer) is removed. 3M HCl (0.4 mL, 10 drops) is added to the ether layer and the vessel is again shaken. The aqueous layer is removed and the ether layer is dried over Na₂SO₄. After 10 minutes, the mixture is filtered through a cotton plugged pipet into dry clean 3mL vial. The Na₂SO₄ is rinsed with ether (0.5 mL X 2). A boiling chip is added and the combined solvent is evaporated with heat until the solution turned cloudy (super saturated).

Purification: Heating is discontinued and crystallization begins. It can be further induced with an ice-bath. The mono-hydrate product is placed in the oven at 100 °C for 1 hour or until the next class period. Note, if the oven is over 100 °C the product will decompose. It is essential that the dried product (no longer a hydrate) be re-crystallized before proceeding to the next experiment.

Spectroscopy: The percent yield is calculated. The melting point and IR spectrum are obtained. (Compare your experimental data (melting point and IR) to that of actual data given for the product. Obtain and interpret an ¹H-NMR, MS and ¹³C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Do not submit your product to the TA, you will use 100 mg of the product acid in a future experiment. If you failed to recover 100 mg, submit a request to your TA for additional starting material so that you will have 100 mg at the beginning of the that experiment.)

Waste Disposal: IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH₂Cl₂ is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na₂SO₄ loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na₂SO₄) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

Mechanism: Students should be able to propose one on their own. It is similar to the first lab. The first equivalent of AlCl₃ is bound to the acid, the other to the ketone carbonyl. On the addition of acid the aluminum complex decomposes.
Experiment 15: Synthesis of 2-methylantraquinone by Friedel-Crafts

Theory & Background: Anthraquinone, is a polycyclic, aromatic hydrocarbon containing two opposite carbonyl groups (C=O) at the 9,10 position. It is a yellow or light gray to gray-green crystal powder that is insoluble in water. In nature, it occurs in plants such as aloe, cascara sagrada, senna, and rhubarb. It is also found in fungi, lichens, and insects as a parent material for the coloring of yellow, orange, red, red-brown, or violet. It is commercially produced by several methods, including oxidation of anthracene with chromic acid, condensation of benzene and phthalic anhydride, followed by dehydration for cyclization, and diene Diels-Alder reaction. Anthraquinone is the most important quinone derivative of anthracene, as it is the parent substance of a large class of dyes and pigments. One of the oldest mordant dye, alizarin, is an anthraquinone derivative. Anthraquinone is a starting material for the production of coloring compounds, antioxidants, and polymerization inhibitors. Its derivatives are widely used as intermediates for dyes, pigments, photographic chemicals, and paints. Anthraquinone is used in paper industry as a catalyst to increase the pulp production yield and to improve the fiber strength through a reduction reaction of cellulose to carboxylic acid. Natural anthraquinones have cathartic properties. Anthraquinones derivatives are also used as drugs. Mitoxantrone, an anti-neoplastic (anti-cancer agent) is but one example.

![Chemical structure of 2-methylanthraquinone](image)

Please regenerate this table in your notebook filling in any of the blanks

<table>
<thead>
<tr>
<th>Reagents</th>
<th>2-(4-toluoyl)-benzoic acid</th>
<th>95% H₂SO₄</th>
<th>2-methylantraquinone</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>equiv</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>MW</td>
<td>240 mg/mmol</td>
<td>98 mg/mmol</td>
<td>222 mg/mmol</td>
</tr>
<tr>
<td>density</td>
<td>solids</td>
<td>1840 mg/mL</td>
<td>solids</td>
</tr>
<tr>
<td>volume</td>
<td>solids</td>
<td>1 mL</td>
<td>solids</td>
</tr>
<tr>
<td>mass</td>
<td>200mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mmol.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>melting point</td>
<td>liquid</td>
<td>175 °C</td>
<td></td>
</tr>
</tbody>
</table>

Procedure: A 5.0 mL vial is charged with 2-(4-toluoyl)-benzoic acid (200 mg). A spin vane and air condenser is affixed to the vial. While stirring at rt, concentrated 18M sulfuric acid (1.0 mL) is added. Caution, concentrated sulfuric acid is highly caustic and corrosive. The vessel is then heated to 100 °C for 1.0 hours.

Work-up: Heating is discontinued and the vial is permitted to cool and a small amount of ice is cautiously added through the throat of the condenser. Caution is observed while adding minute amounts of ice to the hot concentrated sulfuric acid. This process is continued until crystallization stops or the vial volume reaches about 5 mL. The vial should have now reached room temperature. If done slowly enough then the crystals will be large enough to collect by filtration and can be washed with 1.0 mL of concentrated ammonium hydroxide to remove unreacted starting material.
Otherwise

The contents of the vial are cautiously and slowly transferred with to a 25mL erlenmeyer containing 10 eq of solid NaOH so that the acid is quenched and the remaining organic acid starting material goes into the now basic aqueous layer. A complete quench can be confirmed by the addition of Na$_2$CO$_3$; its addition should not emit any CO$_2$ gas if the solution is basic. The quinone can then be extracted with ether, dried over Na$_2$SO$_4$ and concentrated by evaporation.

**Purification:** Recrystallize the product from ethanol or toluene using Norit pellets for decolorization, dry determine the weight and melting point. If off, then the resulting anthraquinone can be further purified by sublimation in a 13 x 100 mm test tube as done in chem 6a. The product should be light yellow in color after sublimation.

**Spectroscopy:** The percent yield is calculated. IR is obtained. Compare your experimental data (IR and melting point) to that of actual data given for 2-methylantraquinone. Obtain and interpret an $^1$H-NMR, MS and $^{13}$C-NMR spectrum from your TA. Include the labeled spectra in your notebook. Submit your entire sample in a disposable tared vial to your TA for scoring.

**Waste Disposal:** IMPORTANT - Syringes do NOT go in the trash. Place syringes in the labeled sharps container! Any residual CH$_2$Cl$_2$ is put in a halogenated organic waste container. Non-halogenated organic waste is put into the non-halogenated waste container. Any silica or Na$_2$SO$_4$ loaded pipets are dumped into solid waste and the glass placed in a sharps container. Free solid waste (silica, Na$_2$SO$_4$) is put into the solids waste container. All aqueous liquids (acetone, ethanol, water) is flushed down the sink after neutralization.

**Mechanism:** Students should be able to propose one on their own mechanism. It is similar to experiment-1.

**Note:** That students are expected to take an exit practical exam which covers all aspects of chem 6b in the next lab period.
We enjoyed having you in Chem 6ab and hope to see you again in Chem 6c where we work on macroscale.

Notebook Submission

Turn your notebook(s) over to your TA before exam week begins for consideration by the instructor.

Chem and Biochem majors

Get to know your organic faculty and find a research job if you are interested.

The organic division strongly endorses the following course electives: Chem 124 (Spectroscopy), Chem 126 (Computational Methods), Chem 127 (Structure and Reactivity), Chem 128 (Organic Reaction Mechanism), Chem 129 (Synthetic Organic Reactions), Chem 199 (Undergraduate research). These courses can be taken in any particular order.

Check in & out procedures

Pick-up the inventory check list from the TA, which you initial along with your TA. Make sure your clean dry plastic bin contains all the equipment listed. If not, get TA approval and go to the storeroom to replace missing items. There should be no other items or chemicals in your bin when you check in & out. If your equipment is not clean, clean it!

Notes from the chem-club

The UCSB Student Affiliates of the American Chemical Society is a group that meets bi-monthly to discuss various topics in Chemistry and Biochemistry. Guest speakers often provide information about research, career opportunities and industry, and other aspects that encompass the field. The group also participates in other activities such as the ACS National Convention, National Chemistry Week Celebration, Outreach, and tours. A barbecue is also hosted at the end of the fall and spring quarters as are trips to the ACS meetings. For more information see:

http://www.chem.ucsb.edu/~chemclub/index.html