### **Experiment 3 – Oxidation of Benzhydrol**

Reading Assignment: <sup>1</sup>H NMR (Mohrig Chapter 22.1-22.7 or McMurry 13.1-3, 13.8-10)
Review Topics: Mohrig Ch 10 (Extraction) & 18 (TLC); McMurry 17.7 (oxidation of alcohols)

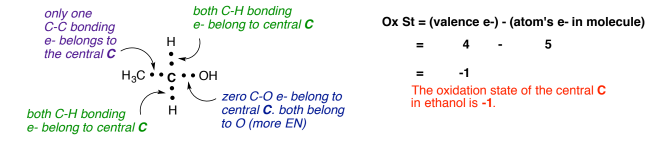
In this experiment, students will perform a simple oxidation reaction of a secondary alcohol. Recall that Oxidation Is a Loss of electrons while Reduction Is a Gain of electrons (OIL RIG). In order to apply this mnemonic, you must know the oxidation states of each atom within the compound, with particular interest on carbon. Carbon can carry oxidation states ranging from -4 to +4. A few examples are shown in **Figure 1** below. It is also common for carbon to carry an oxidation state of -3, -1, +1, and +3.

	Methane	Methanol	Formaldehyde	Formic Acid	Carbon Dioxide
Carbon's	H H- <b>C</b> -H H	H H- <b>C</b> -OH H	O H	O C H OH	O= <b>C</b> =O
Oxidation State	-4	-2	0	+2	+4

Figure 1. Examples of the oxidation levels of carbon

You may have noticed that all of the compounds in **Figure 1** are neutral and carbon has zero formal charge in each example. The concept of oxidation state and formal charge are similar with one important difference in the calculation. Both are calculated by taking the difference between the valence electrons (from the periodic table) and the number of electrons belonging to that atom within the molecule. For a given atom, the valence electrons will never change but the electrons 'belonging' the atom in the molecule will vary depending on number of lone pairs and attachments to more or less electronegative atoms. The important difference in the calculation of oxidation states and formal charge is based on the following assignment of bonding electrons (**Figure 2**). This is how the highlighted carbon in ethanol can have an oxidation state of -1 but a formal charge of zero.

- <u>Oxidation states</u> assign bonding electrons to the more electronegative atom in a bond, except when the two atoms are the same and the bonding electrons are split equally.
- Formal charge splits bonding electrons equally between the two atoms.



Formal Charge, all bonding e- split equally (50.50)...FC = 4 - 4 = 0

The formal charge of the central **C** is ethanol is **zero**.

Figure 2. Calculation of oxidation state and formal charge of the central carbon in ethanol

As stated above, an oxidation reaction is one where an atom loses electrons. In other words, the atom gains a bond to a more electronegative atom (electron hogs!). The examples discussed in this experiment will involve oxygen-containing compounds (alcohols and carbonyl compounds) but there are many other examples of organic oxidation reactions that do not involve oxygen. You should be able to categorize whether the reactions learned in the CHEM 8 series qualify as oxidation or reduction based on the rules outlined above.

**Table 1** below highlights common oxidizing agents and their applications. Selecting the proper oxidizing agent depends on several factors, the most important of which is "does it work?" In choosing the appropriate oxidizing agent, the following issues should be addressed...

- ... Reactivity does it react with the starting material? Is it too reactive or not reactive enough for the desired transformation?
- ... Selectivity will it also react with other functional groups in the molecule?
- ... Ease of use is it toxic and/or does it require special equipment? How is waste handled?
- ... Availability is it commercially available or does it need to be made separately? Is it cost-effective?

Table 1. Common oxidizing agents and applications

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Oxidizing Agent	Main Application(s)	Comments		
Jones Reagent: CrO <sub>3</sub> , H <sub>2</sub> SO <sub>4</sub>	R^OH → R OH	CrO <sub>3</sub> is highly toxic and a carcinogen. High waste disposal cost.		
	OH O R			
Pyridinium chlorochromate (PCC)	$R \curvearrowright OH \longrightarrow R \hookrightarrow H$	Suspected carcinogen, high waste disposal cost.		
Potassium Permanganate (KMnO <sub>4</sub> ) with heat	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Nonselective – many functional groups are oxidized (alkenes, alkynes, alcohols, etc.)		
	$R \longrightarrow CO_2H$	,		
Periodic Acid (HIO <sub>4</sub> )	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	N/A		
Peroxyacids (RCO₃H)	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Common peroxyacid: Meta-chloroperoxybenzoic acid (mCPBA)		
Sodium Hypochlorite (bleach, NaClO)	$R \xrightarrow{OH} R \xrightarrow{R}$	Cheap and easy!		
Dess-Martin Periodinane O O O I –OAc	$ \begin{array}{ccc} OH & O \\ R & & & \\ & & \\ & &$	Easy to use but expensive reagent.		
AcOOAc	R^OH → R H			

In this experiment, the oxidation of a secondary alcohol (benzhydrol) is achieved with commercially available bleach. This reagent is inexpensive and easy to handle with typical personal protective equipment (PPE) including goggles and gloves. Most importantly, it works! However, one issue is presented in using bleach: solubility. Bleach is an aqueous solution of NaClO but many organic compounds, including benzhydrol, are not water-soluble. Thus, a phase transfer catalyst (PTC) is employed to facilitate the reaction.

The mechanism employed by a PTC is similar to that used in soaps. Soaps contain both non-polar and polar (typically ionic) regions so they can absorb grease and also be washed away with water. Quaternary alkylammonium salts such as tetrabutylammonium hydrogen bisulfate ( $Bu_4N^+HSO_4^-$ ) are common examples of PTCs. For the remainder of this discussion, this salt will be abbreviated by  $Q^+X^-$ . These salts are soluble in both water and organic solvents. When  $NaClO_{(aq)}$  is mixed with an immiscible organic solvent such at ethyl acetate (EtOAc), little to none of the NaClO enters the organic phase. However, once a small amount of  $Q^+X^-$  is added, the salts participate in the following equilibrium (eq. 1).

$$Q^+X^- + Na^+CIO^- \longrightarrow Q^+CIO^- + Na^+X^-$$
 (1)

Some of the hypochlorite (CIŌ) ion, the active oxidizing agent, is paired with the tetrabutylammonium cation Q̄<sup>+</sup>. Because Q̄<sup>+</sup> is soluble in organic solvents, it can carry the CIŌ ion from the aqueous to the organic phase where the reaction can occur (**Figure 3**). As the CIŌ reacts in the organic phase, the equilibrium shifts to transport more CIŌ from the aqueous phase to reestablish equilibrium. It is important to note that the salts do not instantly transport from one layer to another. *Vigorous stirring is required to facilitate phase transfer.* This continues until the reaction is complete and the solubility issue is resolved! The applications of PTC are widespread to many other types of reactions, not just oxidations.

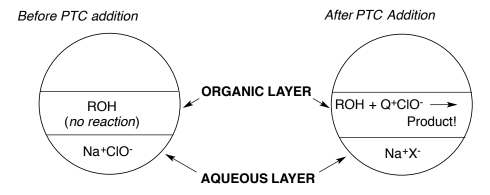


Figure 3. Phase-transfer catalysis (PTC) in an oxidation reaction.

# **Notebook Preparation**

• Purpose – one-sentence description of the purpose in addition to the following scheme:

- Reagent table amount (mg or mL), mmol, equiv\*, MW, bp or mp, density, and one-word
  hazards for each of the chemicals in the scheme above. Leave space in the table to write
  the actual amounts of starting materials used.
  - \* equiv = molar equivalents as defined by the limiting reagent (in this case benzhydrol). After calculating moles of each reagent, indicate the mole ratio of each relative to benzhydrol.
- Procedure hand-written, step-by-step procedure. Reproduce Table 2 and bring the lecture handout for Tables 3 & 4. Include a list of materials.
- Safety & Clean-up copy **Table 5** into your notebook.

## **EXPERIMENTAL PROCEDURE** - Students work individually on this experiment.

Reaction Preparation and Set-up: TLC will be used to monitor reaction progress. Prepare TLC standards and plates before setting up the reaction. Make solutions of the standards (benzhydrol and benzophenone) in small test tubes. This does not require careful measuring, but do be conservative. Dissolve a small amount of the compound (microspatula tip) in ethyl acetate (EtOAc, 1 mL). Obtain three TLC plates, carefully handling by the edges without bending, and gently spot the plate at the origin with a capillary tube (not a melting point capillary). Create one lane for benzhydrol, another for benzophenone, and leave a space for the reaction mixture to be spotted later. Be sure to record which lane is which in your notebook. Take note of the solvent in the TLC chambers.

In a 25-mL Erlenmeyer flask equipped with a magnetic stir bar, add 0.37 g ( $\pm$  0.01 g) of benzhydrol, 5 mL of commercial bleach (approximately 0.7 M NaClO), 5 mL of ethyl acetate (EtOAc), and 40 mg ( $\pm$  5 mg) of tetrabutylammonium hydrogen sulfate (Q $^{\dagger}$ X $^{-}$  or Bu<sub>4</sub>N $^{\dagger}$ HSO<sub>4</sub>). Secure the flask to a ring stand, loosely stopper, and *stir vigorously on a stir plate without heat*.

Monitoring Reaction Progress: After about 10 minutes, stop stirring to allow phase separation and remove a small aliquot of the upper layer of the reaction by touching the tip of a capillary tube to the top of the reaction solvent. Spot the TLC plate with this aliquot using a capillary tube alongside the standards. Run the TLC plate using the chambers provided in the fume hood. Do not remove the chambers from the fume hood! Develop the plate with a UV or fluorescence light after evaporating the solvent from the plate in the fume hood.

If starting material is still present in the reaction, continue stirring for another 10 minutes and take another TLC aliquot. A faint spot for benzhydrol may still appear on a visualized plate, even when the reaction is complete. When there is no *dark* spot for benzhydrol in the reaction mixture, you may consider the reaction to be complete. The 10 minutes is counted from the first aliquot (20 min total). By the time you run the first TLC plate, it's probably time to run the second! Continue taking aliquots at 10-minute intervals until the reaction is complete. If the reaction is taking longer than 40 minutes, stop the reaction and proceed to *Reaction Work-Up*.

\* It is acceptable to obtain 10 mg more or less benzhydrol and 5 mg more of less Q<sup>+</sup>X<sup>-</sup>. Record the exact mass obtained, including the uncertainty of the balance used.

E3-4

Reaction Work-Up: Transfer the completed reaction mixture to a screw-cap test tube and remove the aqueous layer with a pipet. Wash the organic layer with 3 mL of brine (sat. NaCl) followed by a wash with 2 mL of water – mix, invert, then remove the aqueous layer after each portion of brine or water is added. Dry the organic layer over MgSO<sub>4</sub>, gravity filter using a pipet with cotton plug, and collect the filtrate in a pre-weighed 25-mL round-bottom flask (RBF). Concentrate using a rota-vap and weigh the product. Pro-tip: the product rarely crystallizes in the rota-vap bath. When the solvent appears to have evaporated, take the flask off the rota-vap and swirl in the ice bath to crystallize. You can still proceed with the product in liquid form.

Analysis: Obtain the IR of the product. The IR of starting material is online and should also be posted in the instrument room. Record the identifying peaks in your notebook. Sketch the final TLC plate into your notebook and calculate the R<sub>f</sub> values for each spot. Report your data in table format (on the next page). Analyze the <sup>1</sup>H NMR spectra of benzhydrol and benzophenone (provided in lecture) using the following table format. Reproduce Table 2 into your notebook. Tables 3 & 4 are part of the benzhydrol NMR handout from lecture (also available online).

**Table 2.** TLC Results (include solvent here)

Sample	$R_{f}$	Identity	
Benzhydrol			
Benzophenone			
Reaction Mixture (x minutes)		Benzhydrol (if present)	
* add more rows as needed		Benzophenone	

**Table 3.** <sup>1</sup>H NMR Analysis of Benzhydrol (include the structure with H's labeled **A-E**)

	Integration	Expected Chemical Shift	Observed Chemical Shift
Signal	(# of H's)	(ppm)	(ppm)
Α			
В			
С			
D			
E			

**Table 4.** <sup>1</sup>H NMR Analysis of Benzophenone (include the structure with H's labeled **A'-C'**)

Signal	Integration (# of H's)	Expected Chemical Shift (ppm)	Observed Chemical Shift (ppm)
A'			
B'			
C'			

**Table 5.** Clean-up & Safety

Clean-up	Safety		
Liquid waste: aqueous layers and contents of	Ethyl acetate is <i>flammable</i> .		
rota-vap trap			
Solid waste: MgSO <sub>4</sub> , pipets, filter pipets,	Benzophenone, benzhydrol, and Bu <sub>4</sub> NHSO <sub>4</sub>		
capillary tubes, and TLC plates	are irritants.		
After analysis, dispose of your product in the	Sodium hypochlorite is an oxidizer. It will		
liquid waste using a very small amount of	bleach your clothes so consider your wardrobe		
ethanol from a wash bottle to aid the transfer.	for the day!		
Wash all glassware and wipe down counters.	Wear gloves & goggles throughout the		
	experiment.		

#### **Introduction: Pre-Lab Questions**

1. Which atom is oxidized in the reaction of benzhydrol with bleach? Redraw the structures and indicate the oxidation number of that atom in the starting material and product.

- 2. What are the main differences you expect to find between the IR of the starting material and product?
- 3. Briefly explain how phase transfer catalysts work and why one is necessary in this experiment.
- 4. What are some of the advantages of using bleach as an oxidizing agent? What other oxidizing agents could be used to carry out the same transformation?
- 5. What are the two solvents used in the reaction? Will the aqueous layer be on the top or bottom in the extraction?
- 6. Calculate the moles of each reagent used, identify the limiting reagent, and calculate the theoretical yield of benzophenone (recall that catalysts cannot be limiting). Show your work.

## **Results: In-Lab Questions**

- 1. (5 points) Report the yield of product (mg and %). Briefly discuss any parts of the procedure that may have caused the yield to be lower than 100%. Be specific. List the ILE for each instrument used and percent intrinsic error for each measurement.
- 2. (5 points) Report the TLC results (mobile phase, R<sub>f</sub> values, and identification) of TLC analysis in table format (**Table 2**) and explain how you decided to stop the reaction. Briefly explain why the separation was successful by comparing the polarity of the sample and mobile phase.
- 3. (5 points) Compare the IR of the starting material and product. Briefly explain which peaks signify reaction completion, including functional group, bond, and stretching frequency. Attach the IR spectrum of product to the back of the report. No IR table is necessary.
- 4. (10 points) Interpret the <sup>1</sup>H NMR spectra of benzhydrol and benzophenone by re-drawing the structures with labels and re-producing **Tables 3 & 4** into your word processing document. Answer the following in one sentence: **Which NMR peak(s) best distinguish starting material from product?**
- 5. (5 points) Perform a literature search to find a journal article from *Tetrahedron Letters* (*Tet. Lett.*) with "benzophenone" in the title. Web of Science is an excellent search tool and requires on-campus access. Give a one-to-two sentence explanation of the role of benzophenone role in this publication and provide a proper citation to follow your answer (Technical Writing Guidelines, Part E).

### **Experimental Methods**

See Technical Writing Guidelines, Part D for content and format. A sample experimental section is posted online that contains much more detail than 8M students will need to provide! Do include TLC conditions ["reaction progress was monitored by TLC (solvent)"] but do not include NMR data in this section.

Experiment adapted from...

Palleros, D. R. Experimental Organic Chemistry, Wiley: New York, 2000; pp. 255-257.

Exp 3 - Oxidation of Benzhydrol		Name
Section Day	Time	TA Name

# CHEM 8M GRADING RUBRIC - Use as cover page for report

SECTION	INSTRUCTOR COMMENTS	POINTS ASSIGNED
IN-LAB QUIZ		/ 5
LAB REPORT		
ABSTRACT One paragraph, 4-6 sentences: Purpose, procedure, main result(s), and conclusion(s).	NONE	0 / 0
INTRODUCTION Original responses to pre-lab questions with TA initials		/ 30
RESULTS  The main results are stated, as outlined in the in-lab questions, using complete sentences.		/ 30
EXPERIMENTAL METHODS The experimental details (including final amount used and obtained) are briefly described in a few sentences.		/ 20
NOTEBOOK PAGES  Proper format: reaction scheme, chemical info table, procedure, safety, waste, and clean up notes		/ 20
NEATNESS AND ORGANIZATION Proper grammar and format per instructions in syllabus and writing guidelines		/ 10
LAB TECHNIQUE & CLEAN UP Lab space left clean, proper technique, instructions followed, checked in with TA before leaving.		/ 10
	LAB REPORT TOTAL	/ 125