

# Experiment 7

Synthesis and Analysis of  
those same old **Moldy Green Crystals**  
from before spring break

Part 3: Spectrophotometric Determination of Iron Content

CH 204 Spring 2007

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# Two Weeks Ago

## Redox Chemistry

Oxidation — loss of electrons

Reduction — gain of electrons

Balancing redox reactions

Titration with  $\text{KMnO}_4$

# Today's lab in a nutshell

Parts **2** and **3** of the procedure in the lab manual.

- 1) Mix up a series of standards by diluting from a stock solution**
- 2) Measure the absorbance of each of the standards**
- 3) Make a calibration curve by plotting absorbance vs concentration**

# Some quick vocabulary

A **STOCK SOLUTION** is a more concentrated solution that we know the concentration of, and it's the solution we start with when we're going to make a series of standards. The concentration of our iron stock solution is  $0.0188 \text{ g/L Fe}^{2+}$ .

**STANDARDS** are solutions with known concentrations that are around the same concentration as our sample. We will determine our sample concentration by comparing it with our standards.

# Some more quick vocabulary

The **ANALYTE** is the thing we're analyzing for. In this lab the analyte is an orange-colored complex ion  $\text{Fe}(\text{phen})_2^{2+}$ . Our standard solutions and our sample solution contain the same analyte.

Our **SAMPLE** is the solution that we don't know the concentration of, and that's what we're trying to figure out. We'll do this one next week.

An **ALIQOT** is a measured portion of a larger volume.

## Part 2 — make up the standard iron solution

1. Get 10 mL of the iron solution from the hood, and pipette 5 mL into a 25 mL volumetric flask.

That's a 1 to 5 dilution of the original concentration.

2. Add 1 mL of hydroxylamine,  $\text{NH}_2\text{OH}$   
2 mL sodium acetate, and  
8 mL 1,10 phenanthroline
3. Fill the volumetric flask up to the line with deionized water using a dropper pipette, then mix it, cap it off and let it sit for 20 minutes for the reaction to occur.

## Part 3 — Make Individual Standards

1. Get five test tubes and label them 1, 2, 3, 4, 5. Write directly on the glass with your marker.

Using a **graduated pipette**, add that many milliliters of the orange solution that you prepared in Part 2 to each test tube.

Using the graduated pipette again, fill each test tube to 5 mL total by adding 4, 3, 2, 1, and 0 mL of deionized water to test tubes 1-5 respectively.

# A whole lotta dilutin' goin' on!

When we mix up the standards in the test tubes, each one is diluted by a different factor:

1 was diluted 1 to 5

2 was diluted 2 to 5

3 was diluted 3 to 5

4 was diluted 4 to 5

5 was not diluted in this step.



# Calculating final concentrations

To find the final concentration of each of the standards, we have to convert from grams/L to moles/L and then multiply by the dilution factor for each one:

$$\text{Original Concentration (M)} \times \frac{1}{5} \times \text{test tube dilution factor}$$

This dilution was in Part 2

This dilution is in Part 3

$$1: \text{Conc.} \times \frac{1}{5} \times \frac{1}{5}$$

$$2: \text{Conc.} \times \frac{1}{5} \times \frac{2}{5}$$

$$3: \text{Conc.} \times \frac{1}{5} \times \frac{3}{5}$$

$$4: \text{Conc.} \times \frac{1}{5} \times \frac{4}{5}$$

$$5: \text{Conc.} \times \frac{1}{5} \times 1$$

# Spectrophotometry!

Spectrophotometers are the most widely used analytical instruments in the world, except for the analytical balance, and they're about as easy to use as an analytical balance.

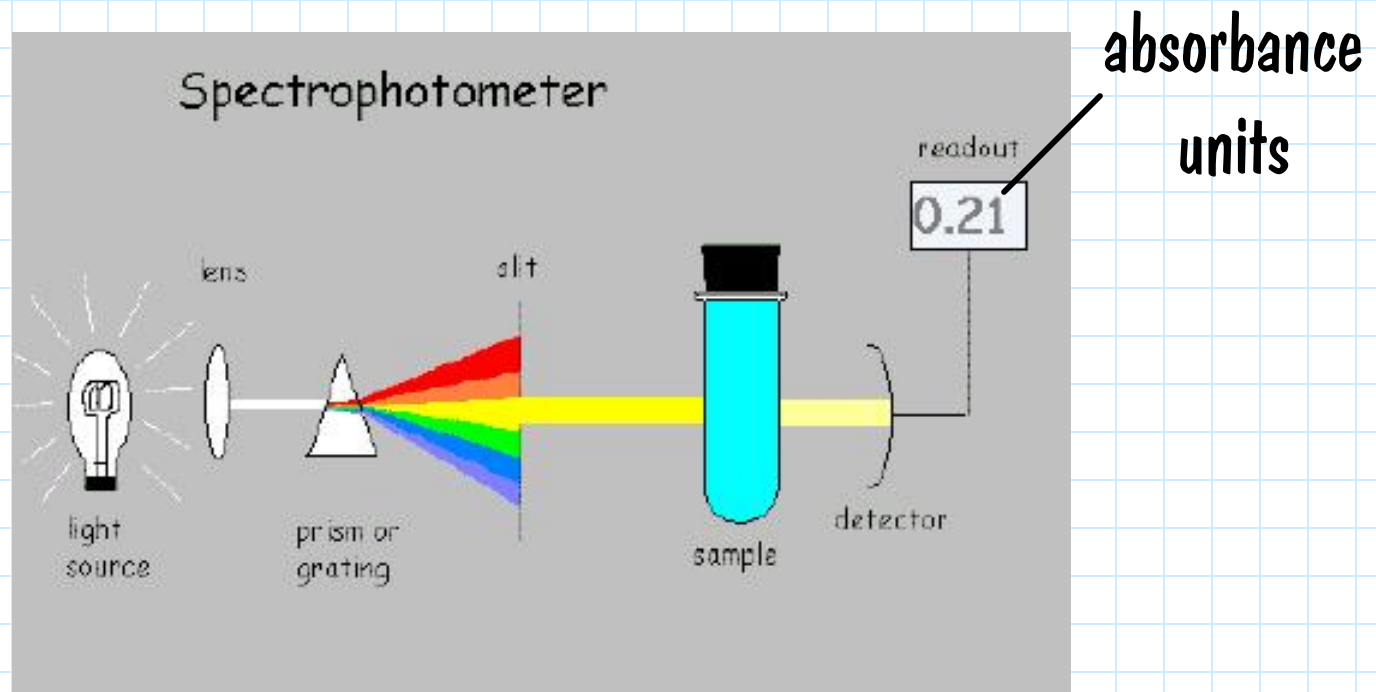
"But what does a spectrophotometer look like?" you are wondering,  
"And how does it work?"

I'm glad you asked!

Looks like this



# Works like this



Anything that is colored has color because it absorbs some wavelength (or wavelengths) of visible light.

# Using the spectrophotometer

Place a **cuvette** full of deionized water into the instrument.  
This is your blank. Press the button that says **0 ABS**.

Remove the blank and put in a cuvette containing your first standard. The display will automatically read out the absorbance. Record this value.

# Lather, Rinse, Repeat

Repeat this procedure for each of your five standards.

Insert the blank before each measurement to make sure the blank reads 0 absorbance units, then insert the next sample.

2 cuvettes to a customer. Reuse the sample cuvette.

# How not to screw up this part

- 1) Rinse the cuvette twice with the solution you are about to measure before you put it in the instrument
- 2) Wipe the outside of the cuvette clean using Kim-Wipes. No fingerprints, no wetness on the outside.
- 3) No bubbles in the solution.
- 4) Fill the cuvettes at least  $3/4$  of the way up.

But what do these absorbance values tell us?

# Beer's Law

Beer's Law says that **absorbance** depends on three factors: molar absorptivity, concentration, and path length.

$$A = \epsilon cl$$

Sometimes written as  $A = \epsilon bc$

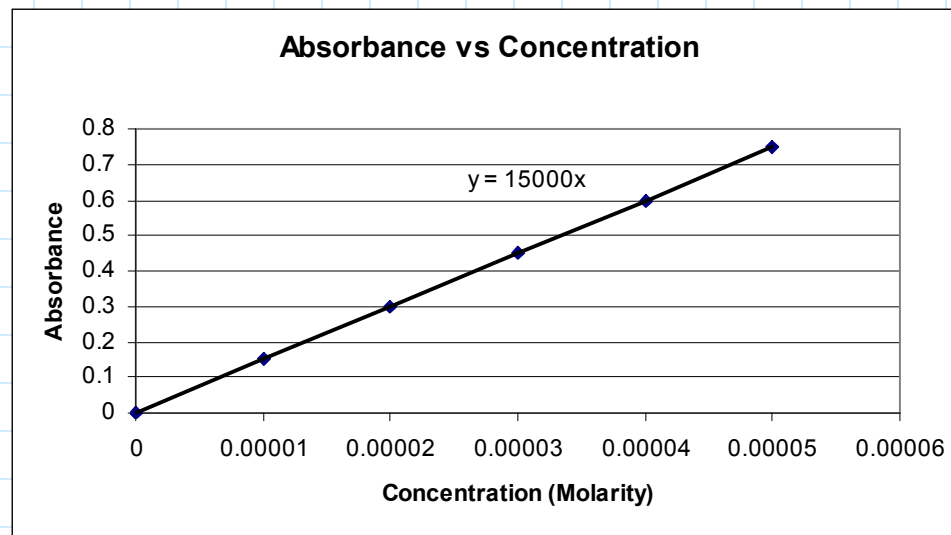
or

$$A = abc$$



# Beer's Law plots

When we plot Absorbance versus Concentration, the slope of the line is equal to  $\epsilon l$ . In our case  $l=1$ , so the slope of the line is equal to the molar absorptivity for  $\text{Fe(phen)}_3^{2+}$ .



## After you have your data

Enter the absorbance and concentration values into Excel.

Plot Absorbance (y-axis) versus concentration (x-axis).

Include 0,0 as a data point — that is your blank.

You should get a straight line, and the slope of the line is your molar absorptivity,  $\epsilon$ , in units of  $M^{-1}cm^{-1}$ .

# Next week

Parts 1 and 4 of Experiment 7 (analyze the sample)

No lab report this week!

Turn in Post-lab 7 at the beginning of lab next week

Quiz next week covering Experiment 7