

## Experiment 7b

Still More Analysis of  
those **Stupid Green Crystals** We Made  
Like a Million Years Ago in Lab

Part 3: Spectrophotometric Determination of Iron Content

CH 204 Spring 2008

Dr. Brian Anderson

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## Last Week

Made standard solutions from a stock solution

Fun with dilutions!

Spectrophotometry — Beer's Law

## 'Tis the season...

Throw a pair of sweat pants in your lab drawer if you're going to be wearing shorts around campus in warmer weather.

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## This week

Convert our **green crystals** into an **orange solution**,  $\text{Fe}(\text{phen})_3^{2+}$ , then use a spectrophotometer to measure how much light this solution absorbs at 510 nm.

Use the molar absorptivity determined from the calibration curve to calculate how much  $\text{Fe}(\text{phen})_3^{2+}$  is in our final sample solution.

Back-calculate through the sample dilutions to determine how much Fe was in the original sample.

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## Lab Procedure, Part 1

1. Weigh out 0.15 g of green crystals and dissolve in deionized H<sub>2</sub>O. (Do this right there in the weighing boat.) Transfer the dissolved sample to a 25 mL volumetric flask.
2. Add 8 mL of 6 M H<sub>2</sub>SO<sub>4</sub>, and fill to the line with deionized water using a disposable pipette.

Your sample is now dissolved in 25 mL of solution.

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## Part 1, continued...

4. Pipette 5 mL into a 30 mL beaker, add about 10 mL deionized water, heat and stir. Start the reaction right away!
5. Add KMnO<sub>4</sub> dropwise until the solution turns pink.  
(Around 30-50 drops.) Yellow → colorless → pink.
- 6-7. Transfer the solution to a 25 mL volumetric flask and let it cool on the dark part of the lab bench.
8. There is no step eight.
9. Fill the volumetric flask to the mark using deionized water.

The sample has now been diluted 1 to 5 from the original concentration.

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## On to Part 4

1. Pipette 5 mL of your sample from part 1.9 into a 25 mL volumetric flask.

The sample has now been diluted 1 to 5 TWICE, or 1 to 25.

Add 1 mL of hydroxylamine, NH<sub>2</sub>OH

2 mL sodium acetate, and

8 mL 1,10 phenanthroline

Fill to the mark with **PHENANTHROLINE!**

Let it sit for 20 minutes.

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### Remember...

...FILL THE ORANGE SOLUTION TO THE MARK  
WITH **PHENANTHROLINE!**

Not with water.

¡No con agua, amigo!

Nicht mit der wasser.

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### Measure the absorbance

You want an absorbance reading that is between your lowest  
and highest standard from last week.

If it's too high, do another dilution and test it again.

**RECORD ALL DILUTIONS!**

**RECORD ALL MEASURED ABSORBANCE VALUES!**

Make all of your dilutions **5 ml** to **25 ml**.

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### "Triplicate" analysis

Once you've got an absorbance reading within the  
range of your standards, dump the cuvette, rinse it  
twice with your solution, and make a second  
absorbance measurement. Then do this again and  
make a third absorbance measurement. Record all  
three absorbances in the table on page 59.

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### Don't forget about those dilutions

When you determine the concentration of your sample from the graph, remember that you have to back-calculate through all the dilutions you made in order to figure out the original concentration you started with.

Your total dilution factor will be  $5^x$  where  $x$  is the number of dilutions you made.

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### Report buried in the procedure

In Part 4 of the experimental procedure, steps 4 – 9 are actually part of the REPORT, not part of the experiment.

These calculations are covered on the cheat sheet from Dr. Leytner.

Make sure you do them!

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### Looking ahead

- The final three labs (Thermochemistry, Acid-Base Equilibria, and Kinetics) will be done in pairs.
- Pre-Lab 8 is longer than previous pre-labs.
- Check full of calculations that you will need in order to do the lab write-up.
- Start on this EARLY

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## Quiz 7 of 9...

Get all the post-lab problems correct from now on.

Start on the post-lab the day you do the lab. Get help from the TA's or from me.

If you understand how to do the post-labs, you'll do better on the quizzes.

No quiz next week.

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