## CHEM 133 Final Exam May 18, 2017 167 Points

Name \_\_\_\_\_

h = Planck's constant =  $6.626 \times 10^{-34} \text{ J} \cdot \text{s}$ , speed of light in a vacuum = c =  $2.998 \times 10^8 \text{ m/s}$ Boltzman Constant = k =  $1.38 \times 10^{-23} \text{ J/K}$ , Avogadro's number = N<sub>A</sub> =  $6.02 \times 10^{23}$  items/mol F = Faraday's constant =  $96,500 \text{ C/mol e}^-$ Pascal's triangle: 1 1 1 2 1 1 3 3 1

SHORT ANSWER SECTION: (Each question worth 4 points)

Method A has a limit of detection of 20 μg while method B has a limit of detection of 300 ng (nanograms). Which of the following is the best statement in comparing method A to method B?
 a) method B is more accurate b) method B is more precise
 c) method B is more sensitive d) method A has a greater range of response

2. A fluoride selective electrode has a useful output range of 0.6 to 2.1 V and is being recorded by a digitizer with an input voltage range of 0 to 1 V. Noise is not a major issue. What type of analog signal processing is needed?

a) RC high pass filter b) RC low pass filter c) a voltage divider d) a voltage amplifier

3. If the reference electrode is acting as the cathode and the analyte electrode as the anode, the cell voltage = (where  $E_{ref}$  and  $E_{analyte}$  refer to the reference and analyte electrode potentials, respectively): a)  $E_{ref}$  b)  $E_{analyte}$  c)  $E_{analyte} - E_{ref}$  d)  $E_{ref} - E_{analyte}$ (assume  $E_{cell} > 0$ )

4. Which of the following monochromator components is designed to disperse light?a) collimating opticsb) gratingc) focusing opticsd) exit slit

5. In NMR, the light transducer is:a) an antenna to pick up radio frequency wavesb) tc) the Fourier Tranformerd) t

b) the magnet shimsd) the permanent magnet

6. List a main component (one of 3) of a mass spectrometer and a specific example of this component (e.g. if this question were asking about UV-visible spectrometers you could say <u>lamp</u> and <u>deuterium lamp</u>). Main component \_1) ion source, 2) mass analyzer, 3) ion detector\_\_\_\_\_ Specific Example: \_1) EI, CI, ESI, MALDI, DESI, APCI 2) quadrupole, time-of-flight, ion trap, etc., 3) Faraday cup, electron multiplier, MCP\_\_\_

7. It is desired to remove free fatty acids from biodiesel. These compounds are only weakly soluble in water if in their molecular form, but can be dissolved if fatty acids exist as ions. A way to remove these from biodiesel is to wash the biodiesel with water containing:

a) an acidic buffer b) a basic buffer c) dissolved NaCl

8. Which of the following chromatograph block diagrams has components in the correct order? a)



9. GC is desired to separate two compounds. In looking up compound properties, a scientist finds both compounds have the same boiling point temperature but different polarities (e.g. octanol – water partition coefficients). He will expect the resulting chromatogram:
a) to show slower elution of the compound with polarity similar to the stationary phase

b) to show faster elution of the compound with polarity similar to the stationary phase c) to always give overlapping peaks

d) to give unexpected results because he was supposed to look up the melting point temperature

10. A separation is performed using normal phase HPLC with a mobile phase of 90% hexane/10% 2-propanol and a 150 mm x 4.6 mm silica column. Because solute peaks are found to elute quickly and overlap, it is desired to increase retention. This can be accomplished by:

a) using 95% hexane/5% 2-propanol
b) using a 150 mm x 8.0 mm column
d) it is impossible to increase retention in this case

11. The primary reason going to smaller diameter open tubular columns and thinner stationary phase films in GC results in smaller H values under the same linear velocities is:a) the flow rate will be lowerb) molecular diffusion is decreased

c) mass transport to and within the stationary phase is faster

d) the variations in paths around the smaller particles is greater

12. In GC analysis, broad or tailing peaks often occur and give poor performance. If it is expected that this is from column overload, an approach in reducing this is to:

a) concentrate the samples before injection b) make sure splitless injection is being used

c) decrease the fraction being injected by split injection

d) use a smaller diameter column

## SECTION II. PROBLEM SECTION. Show work - use the back side if needed

1. A student is building a tube heater by wrapping heating wire around a tube. The heating wire has  $0.60 \text{ }\Omega/\text{m}$ . See the figure below.



a) Assuming the heating wire is the only source of resistance in the circuit, what is the current needed in the heating wire if it is desired to to provide 35 W to the tube heater? (6 pts)

P = IV = 35W and V = 10.0V, so I = P/V = 3.5A

b) What length of wire would be needed to produce the desired 35 W of heat? (6 pts)

 $V = IR \text{ or } R = V/I = 10V/3.5 \ \Omega = 2.86 \ \Omega \text{ or } L = 2.86 /(0.60 \ \Omega/m) = 4.8 \ m$ 

2. A researcher is developing an electrode to measure oxalate  $(C_2O_4^{2-})$  in water. She needs to determine the standard reduction potential  $(E^\circ)$  for the reduction reaction:

1)  $CdC_2O_4(s) + 2e^- \leftrightarrow Cd(s) + C_2O_4^{2-}$ Given that E° for 2)  $Cd^{2+} + 2e^- \leftrightarrow Cd(s)$  is -0.402 V and K<sub>sp</sub> for 3)  $CdC_2O_4(s) \leftrightarrow Cd^{2+} + C_2O_4^{2-}$  is 8.72 x 10<sup>-5</sup>, calculate the reduction potential for the first reaction. Note: standard conditions include 25°C. (12 pts) *Method A: rxn 1* = *rxn 2* + *rxn 3* so  $\Delta G_1^{\circ} = \Delta G_2^{\circ} + \Delta G_3^{\circ}$   $\Delta G_1^{\circ} = -nFE_2^{\circ} + -RTlnK = -2(96500)(-0.402 V) - (8.314)(298)ln(8.72 x 10^{-5})$   $\Delta G_1^{\circ} = 77,600 J/mol + 23,200 J/mol = 100,700 J/mol = -nFE_1^{\circ} or$  $E_1^{\circ} = -100,700/[(2)(96500)] = -0.522 V$ 

*Method B:* We can determine  $[C_2O_4^{2-}]$  in reaction 1 that is in equilibrium with  $[Cd^{2+}]$  in reaction 2 and then use the Nernst Equation.

 $E_2 = E_2^{\circ} - 0.05916/2\log Q$  where  $Q = 1/[Cd^{2+}]$  and  $[Cd^{2+}] = K_{sp}/[C_2O_4^{2-}]$ and  $[C_2O_4^{2-}] = 1$  M (standard state) so  $E_2 = E_2^{\circ} - 0.05916/2\log(1/8.72 \times 10^{-5}) = -0.522$  V

3. Some infrared detectors measure temperature increases (e.g. due to infrared light hitting a thermistor). If the minimum temperature increase detectable by a thermistor corresponds to a heat flux of  $1.0 \times 10^{-6}$  J (over a 1 s period), calculate the minimum number of photons needed for detection for infrared radiation with a wavenumber of 800 cm<sup>-1</sup>. (10 pts)

 $E = hc/\lambda \text{ and } \lambda = 1/\breve{v} \text{ or } E = hc\breve{v} = (6.626 \text{ x } 10^{-34} \text{ J} \cdot \text{s})(2.998 \text{ x } 10^8 \text{ m/s})(800 \text{ cm}^{-1})(100 \text{ cm/m})$  $E = 1.59 \text{ x } 10^{-20} \text{ J/photon}$ 

No. photons =  $(1.0 \times 10^{-6} J)/(1.59 \times 10^{-20} J/photon) = 6 \times 10^{13} photon$ 

4. In flame atomic absorption spectroscopy a number of processes can decrease the atomization efficiency. (10 pts)

a) Give an example of one process that decreases atomization efficiency

1) formation of non-volatile salts

2) ionization

3) oxide/hydroxide formation

b) Give a method to reduce the effect of this process
1) use hotter flame/add compound to bind to anions/standard addition
2) use compound that readily ionizes/standard addition
3) use fuel rich flame/standard addition (only if matrix affects this)

5. Alkyne protons (see figure below) are influenced by electron circulation in the two pi bonds resulting in an induced magnetic field as shown below. Explain how you would expect this to affect the chemical shift (e.g. upfield or downfield) of the HC $\equiv$  protons (compared to protons with the same electron density from the HC bond). (6 pts)



We can see that  $H_{induced}$  is in opposite direction from  $H_{applied}$ . This means  $H_{nucleus}$ =  $H_{applied}$  -  $H_{induced}$ . Because  $H_{nucleus}$  is less than  $H_{applied}$ , we would need a larger (**upfield**)  $H_{applied}$  value to obtain the same resonance frequency as a proton with the same electron density.

6. A compound is analyzed by GC-MS (using electron impact ionization) and is suspected to be dichlorobenzene ( $C_6H_4Cl_2$ ). Given information below,

Element/Isotope	Mass (amu)	Abundance (relative to main isotope)
$^{1}\mathrm{H}$	1.01	100
$^{12}C$	12.00	100
<sup>35</sup> Cl	34.97	100
<sup>37</sup> Cl	36.97	32.0

a) predict the mass to charge ratio of three mass peaks associated with the parent ion (main peak plus isotope peaks only from different Cl isotopes). Assume no multiple charges. (6 pts) main ion mass = 6(12) + 4(1.01) + 2(34.97) = 145.98 amu = m/z (no loss or gain of H with EI)  $1^{37}Cl - mass = 145.98 - 34.97 + 36.97 = 147.98$  amu and  $2^{37}Cls$ : 149.98 amu

6 - cont. b) Calculate the expected ratio of the abundances of the two isotope peaks in a) relative to the main isotope peak. (10 pts)

147.98 peak = M + 2 peak: M + 2/M abundance = 2(32)/100 (2 because either Cl can be 37) M + 2/M abundance = 64/100 149.98 peak = M + 4 peak: M + 4/M abundance =  $(32)^2/100^2 = (1024/100)/100$  (^2 because both Cls must be 37) M + 4/M abundance = 10.2/100

6 - cont c) The sensitivity of the mass spectrometer can be changed by changing the amount of time the quadrupole spends on specific m/z ion (e.g. by changing the mass range scanned). If the mass spectrometer spent 5.0 ms (milliseconds) on each m/z during the scan and the signal to noise of the main isotope peak was 73, what time is needed to obtain a signal to noise ratio of 10 for the smallest isotope peak? (5 point bonus)

S/N(M + 4 peak) = (0.102)(73) = 7.4 $(S/N)_{desired}/(S/N)_{5ms} = (t_{desired}/t_{5ms})^{0.5} \text{ or } 10/7.4 = (t_{desired}/5 ms)^{0.5} \text{ or } t_{desired}/5 ms = (10/7.4)^2 = 1.83 \text{ or } t_{desired} = 9.0 \text{ ms}$ 

7. A sulfated maltose molecule is prepared and isolated. It is believed that the compound consists of three sulfate groups in which  $-OSO_3H$  replaces -OH. It is analyzed by negative ion electrospray ionization (ESI) – mass spectrometry. The formula for (unsulfated) maltose is  $C_{12}H_{22}O_{11}$ . If it is known that sulfated maltose loses one, two or three H<sup>+</sup>s, predict the m/z value for the three peaks corresponding to these H<sup>+</sup> losses. (12 pts)

Isotope	Mass (Da)
<sup>1</sup> H	1.0078
$^{12}\mathrm{C}$	12.0000
<sup>16</sup> O	15.9949
<sup>32</sup> S	31.9721

Formula for trisulfated maltose:  $C_{12}H_{22}O_{11} + 3SO_3$  or  $C_{12}H_{22}O_{20}S_3$ 

*Molecular mass* = 12\*12.00 + 22\*1.008 + 20\*15.99 + 3\*31.97 = 581.89 = M

*Mass of 3 ions* = M - nH; *so for* n = 1, 2, *and 3*, M - nH = 580.88, 579.87, *and* 578.86 (*ignoring the mass of electrons*)

m/z of ions = M - nH/n; so for n = 1, 2, and 3, M - nH/n = 580.88 Da, 579.87/2 = 289.94 Da, and 578.86/3 = 192.95 Da

8. Two columns, a non-polar DB-1 column and a moderately polar DB-225 column, were tested to separate methyl stearate – MS or  $CH_3O_2C(CH_2)_{16}CH_3$  from methyl oleate – MO or  $CH_3O_2C(CH_2)_7CH=CH(CH_2)_7CH_3$  (cis form) using GC in two trial chromatograms with retention times and peak widths listed below. The columns had similar dimensions (length, diameter and film thickness) and were run using the same temperature program.

Peak	DB-1 Retention	DB-1 Peak width	DB-225	DB-225 Peak
	Time (min)	at base (min)	<b>Retention</b> Time	width at base
			(min)	(min)
CH <sub>4</sub>	1.71	-	1.83	-
(unretained)				
MO	8.53	0.185	10.31	0.213
MS	8.79	0.190	8.84	0.186

a) Determine the retention factor (k) for MO on the DB-1 column. (5 pts)  $k = (t_r - t_m)/t_m = (8.79 - 1.71)/1.71 = 3.99$ 

b) Determine the plate number (N) for the DB-225 column using MO. (5 pts)  $N = 16(t_r/w)^2 = 16(10.31/.213)^2 = 37,500$ 

c) Based on their separations, which compound (MS or MO) is more polar? Explain your answer. (5 pts)

MO – it comes out later on a more polar column

d) Calculate the resolution on the DB-1 column. Are the peaks baseline resolved? (6 pts)  $R_s = \Delta t / w_{ave} = (8.79 - 8.53) / [0.5(0.190 + 0.185)] = 1.39 < 1.5$  so not baseline resolved

e) **Select a column**, which when optimized for a fast separation, should allow better analysis of biodiesel assuming MO and MS are the main biodiesel constituents of interest. Assume that both columns could be run hotter or cooler than the test conditions used above and that there will be no change in  $\alpha$  value with change in temperature. Explain your choice (you may want to perform calculations to justify your selection). Also, **indicate what change in conditions** you would use to optimize the separation. (6 pts)

*DB-225* looks better (resolution is much higher 7.37 vs. 1.39 under start conditions). Since resolution >> 1.5, one can **increase temperature** to cut the time.

9. Additional Problem 3.1 The table below shows the retention times and peak widths for 3 compounds separated by reversed-phase HPLC using a 250 mm length x 4.6 mm inside diameter column with an eluent of 45% acetonitrile, 55% water and a flow rate of 1.00 mL/min:

Compound	Retention time (min.)	Peak Width at half height (min.)	pKa
unretained	1.58	NA	NA
phenol	4.72	0.21	9.98
2-nitrophenol	5.68	0.25	8.39
2-methylphenol	5.91	0.26	10.09

Note:  $w_b = 1.70 * w_{1/2}$ 

a) Based on the elution order, which compound is most polar? Explain. (4 pts) **Phenol** – most polar elutes first on reversed phase (less like stationary phase)

b) Suggest a change in mobile phase without changing pH to increase retention. (4 pts) *Increase %H<sub>2</sub>O* (*the weaker eluent*)

c) If the pH of the eluent is increased to 8.0 (from an assumed pH of ~7 in the first eluent) and the composition remains 45% acetonitrile, 55% aqueous buffer, what will happen to each retention time? (6 pts)

Although this pH is less than any of the pK<sub>a</sub> values, it is close enough to 8.39 (2-nitrophenol), that some of 2-nitrophenol will be ionized. When compounds are ionized, they don't spend any time in the stationary phase. This will cause all compounds to elute sooner. However, because the other two phenols have  $pK_a \sim 10$ , only  $\sim 1\%$  will be ionized and the change to elution will be very minor. So the net effect will be an earlier elution time for 2-nitrophenol and almost no change for the other two phenols.