Chemistry 271 – 23XX
Prof. Jason Kahn
University of Maryland, College Park
General Chemistry and Energetics
Final Exam (200 points total)

Your Name: __________________________________________
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Your SID #: ____________________________________________
General Chemistry and Energetics
Final Exam (200 points total)

Your Section # or time: ________________________________

December 15, 2016

You have 120 minutes for this exam.

Explanations should be concise and clear. There is extra space on the last page if you need it.

You will need a calculator for this exam. No other study aids or materials are permitted.

Generous partial credit will be given, i.e., if you don’t know, guess.

Useful Equations:

\[ K_a = [H^+][A^-]/[HA] \]

\[ K_w = [H^+][OH^-] \]

\[ R = 0.08206 \text{ L} \cdot \text{atm/mole K} \]

\[ k_B = 1.38 \times 10^{-23} \text{ J/K} \]

\[ \Delta S = q/T \]

\[ S = k_B \ln W \]

\[ pH = -\log([H^+]) \]

\[ pH = pK_a + \log[A^-]/[HA] \]

\[ \Delta G^\circ = -RT \ln K_{eq} \]

\[ \Delta G = \Delta H - T \Delta S \]

\[ \Delta G = \Delta G^\circ + RT \ln Q \]

Chemical standard state: 1 M solutes, pure liquids, 1 atm gases

Biochemical standard state: pH 7, all species in the ionic form found at pH 7

\[ ^\circ C = ^\circ K - 273.15 \]

\[ P(v)dv = Cv^2 \exp(-mv^2/2kT) \]

\[ 2.303RT/\mathcal{F} = 0.0592 \text{ Volts at 25 } ^\circ \text{C} \]

\[ \Delta G^\circ = -n\mathcal{F}E_{cell}^\circ \]

\[ \ln k = (-E_a/RT) + \ln A \]

\[ [A] = [A]_0 - kt \]

\[ \ln[A] = \ln[A]_0 - kt \]

Standard hydrogen electrode: \[ 2 \text{ H}^+ (aq, 1 \text{ M}) + 2 e^- \rightarrow \text{H}_2 (g) \]

\[ E_{cell}^\circ = 0.000 \text{ V} \]

Honor Pledge: At the end of the examination time, please write out the following sentence and sign it, or talk to me about it:

“I pledge on my honor that I have not given or received any unauthorized assistance on this examination.”
1. **Fundamental thermodynamics (45 pts)**

(a; 15 pts) Fill in the blanks: As a chemical reaction \( \text{A(g)} \rightleftharpoons \text{B(g)} \), starting with pure A, proceeds from left to __________________ at constant \( T \) and \( P \), \( Q = \) ________________[enter expression for the reaction quotient] increases until \( Q = \) ________________ and then remains constant. At this point we have reached the state of ________________. The change in the state function ________________, denoted \( \Delta \)_______, is zero for converting a small amount of reactants to products or vice versa. This tells us that at these particular partial pressures of A and B, the _______________ of A is ________________ the molar free energy of B.

(b; 25 pts) Multiple T/F: Circle ALL correct answers.

<table>
<thead>
<tr>
<th>Question</th>
<th>Answer 1</th>
<th>Answer 2</th>
<th>Answer 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free energy ( G )...</td>
<td>Is always positive</td>
<td>Is a state function</td>
<td>Is conserved, like ( E )</td>
</tr>
<tr>
<td>When ( G ) is at a minimum...</td>
<td>The system is at equilibrium</td>
<td>( T = 0 ) K</td>
<td>The enthalpy of the universe is a maximum</td>
</tr>
<tr>
<td>( \Delta G ) is useful because...</td>
<td>It can easily be calculated from first principles</td>
<td>It lets us focus only on the system and not the surroundings</td>
<td>It gives us the temperature dependence of ( K_{eq} )</td>
</tr>
<tr>
<td>An exothermic reaction in system that is in a bath that maintains constant ( T ) and ( P )...</td>
<td>Increases the entropy of the surroundings (the bath)</td>
<td>Must take heat in from the bath</td>
<td>Is always spontaneous</td>
</tr>
<tr>
<td>The standard state free energy change ( \Delta G^\circ )...</td>
<td>Depends on temperature</td>
<td>Is always = 0 at equilibrium</td>
<td>Is independent of the actual concentrations of reactants and products</td>
</tr>
<tr>
<td>The actual measured free energy change ( \Delta G )...</td>
<td>Depends on temperature</td>
<td>Is always = 0 at equilibrium</td>
<td>Is independent of the actual concentrations of reactants and products</td>
</tr>
<tr>
<td>The entropy ( S )...</td>
<td>Cannot be measured; only ( \Delta S ) can be measured</td>
<td>( S = 0 ) for a perfect crystal at 0 K</td>
<td>Is decreasing for the universe as a whole</td>
</tr>
<tr>
<td>Spontaneous protein folding is an example of ...</td>
<td>Endothermic ordering</td>
<td>Exothermic disordering</td>
<td>Endothermic disordering</td>
</tr>
<tr>
<td>Time’s arrow has a direction because of the ...</td>
<td>Second Law of Thermodynamics</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Score for the page___________
(c; 5 pts) If you memorized the equation \( G = G^\circ + RT \ln P \), but forgot the sign, how could you figure out whether + or − is logically correct?

2. **Electrochemistry (40 pts)**

Consider a \( \text{Zn}(s)/\text{Zn}^{2+}(aq) \) half cell set up by one of your lab partners, hooked up to what your other lab partner told you is a Standard Hydrogen Electrode (SHE).

(a; 12 pts) Sketch the setup, including labeling the anode, cathode, and salt bridge and indicating the direction of electron flow through the external circuit.

Based on the standard reduction potential tables, the voltage should be +0.76 V for the oxidation of Zn to reduce \( H^+ \) to \( H_2 \).

(b; 8 pts) Write down each half-reaction and the balanced overall redox reaction.
Instead of 0.76V, you measure 0.71 V. You suspect that one, but only one, of the aqueous components was prepared incorrectly, i.e. that either the Zn\(^{+2}\) concentration or the H\(^+\) concentration is not at the standard state. [We ignore any possible overpotential or conductivity complications.]

(c; 8 pts) Calculate the non-standard Zn\(^{+2}\) concentration that would be required for you to observe 0.71 V, assuming that the SHE was set up correctly.

(d; 6 pts) Calculate the pH of the non-standard hydrogen electrode (denoted the $%\$@#*&%$-NSHE) that would be required to give 0.71 V, assuming that the Zn(s)/Zn\(^{+2}\)(aq) half-cell was set up correctly.

(e; 6 pts) Which half-cell is the likely culprit: explain your reasoning?
3. van’t Hoff and DNA (40 pts)

(a; 7 pts) We know that \( \Delta G = \Delta H - T\Delta S \) in general, and \( \Delta G^\circ = -RT \ln K_{eq} \) for chemical equilibrium. Use these two equations to derive the van’t Hoff equation, and indicate the dependent and independent variables.

(b; 6 pts) For DNA hybridization \( W + C \rightleftharpoons W\cdot C \), \( \Delta H^\circ \) and \( \Delta S^\circ \) are both negative: explain why, physically.

(c; 11 pts) The sketch below shows the primary data for a melting curve for a DNA oligonucleotide. Label the axes, indicate the part of the curve where the DNA is almost all double stranded (ds), indicate the part where it is single-stranded (ss), and estimate the “melting temperature” Tm, the temperature at which half the DNA is in single-stranded form and half is double stranded.

Score for the page
The value of $T_m$ is given by the following equation: 

$$T_m = \frac{\Delta H^\circ}{\Delta S^\circ + RT \ln(C_T)}$$

where $C_T$ is the total input strand concentration of the two input single-stranded DNAs (W and C), and $R = 1.987 \text{ cal/mole K}$ in the traditional non-SI units used in this field. In this example, $\Delta H^\circ = -71.3 \text{ kcal/mole}$, $\Delta S^\circ = -194 \text{ cal/mol K}$, and $C_T = 6 \mu\text{M}$.

(d; 8 pts) Without doing a numerical calculation, use the equation to say whether $T_m$ increases or decreases as $C_T$ increases, and give a rationale for this trend based on LeChatelier or kinetic arguments.

(e; 8 pts) Finally, calculate the $T_m$ and sketch the melting curve you would get if you melted a different oligonucleotide, with $\Delta H^\circ = -73.0 \text{ kcal/mole}$ and $\Delta S^\circ = -194 \text{ cal/mol K}$, at 6 $\mu\text{M}$ $C_T$ as above.
4. Environmental thermodynamics, e-chem II (45 pts)

(a; 5 pts) What does the “350” in www.350.org mean?

(b; 8 pts) Where does the energy released in combustion come from? In terms of redox, explain why burning natural gas provides about twice as more free energy per carbon atom released than burning coal or wood.

(c; 6 pts) List one lifestyle decision (not necessarily your own) and one society-level choice that could lead to a moderation of the rate of increase in atmospheric CO\(_2\).

(d; 3 pts) List one greenhouse gas besides CO\(_2\).

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(e; 9 pts) The job of PSII is to make a great oxidizing agent (the P680$^+$ cation). What does it oxidize? The job of PSI is to make a strong reducing agent. Where do the electrons from this reducing agent go in the end? Why is it necessary to use two photons in the overall process instead of just one?

The movie “The Martian” depends on a machine, based on an actual proposed NASA device code-named “MOXIE,” that somehow converts CO$_2$ collected from the Martian atmosphere to O$_2$ for Matt Damon to breathe. MOXIE expels carbon expelled as CO. We will oversimplify the chemistry by imagining that it occurs in aqueous solution. (In the real future MOXIE will operate without H; if Mars had unlimited water, we would get oxygen simply by electrolysis of water. And if wishes were horses, beggars would ride.)

(f; 4 pts) Balance the half-reaction for CO$_2$ reduction to CO in acidic solution, for which $E^{\circ}_{\text{red}} = -0.53$ V.

(g; 4 pts) Balance the half-reaction for the oxidation of water to produce O$_2$ in acidic solution, for which $E^{\circ}_{\text{ox}} = -1.23$ V.

Score for the page___________
(h; 6 pts) Write down the overall balanced electrochemical reaction for MOXIE and calculate the standard state cell voltage. Is this voltage pH-dependent? Why or why not? Circle one: this is an electrolytic cell or a galvanic cell?

5. **Statistical thermodynamics (30 pts)**

Consider distributing 6 quanta of energy among 8 particles. The two configurations comprising the largest numbers of microstates in the ensemble comprise about half of the total number of microstates.

(a; 7 pts) One of two dominant configurations has two particles with two quanta each and two particles with one quantum each. Draw two microstates that belong to this configuration, and calculate how many microstates are in it.
(b; 7 pts) The other configuration in this ensemble has particles occupying as many different energy levels as possible (i.e. particles that have as many different numbers of quanta as possible). Draw a microstate in that configuration and calculate how many microstates there are in it.

(c; 12 pts) Whose distribution comes from adding up the overall number of particles with each energy? Sketch the distribution. What is the relationship between entropy and microstates? Why do we use the logarithm?
(d; 4 pts) We know that the number of microstates in a configuration is maximized if every particle in has a different energy – why didn’t a configuration with every particle at a different level show up as the winner?