last name			first name							sign	signature						
1 1 H 1.008	2											13	14	15	16	17	18 2 He
3 Li 6.941	4 Be _{9.012}											5 B 10.81	6 C 12.01	7 N 14.01	8 O 16.00	9 F 19.00	10 Ne 20.18
11 Na 22.99	12 Mg _{24.31}	3	4	5	6	7	8	9	10	11	12	13 Al _{26.98}	14 Si _{28.09}	15 P 30.97	16 S 32.07	17 Cl 35.45	18 Ar 39.95
19 K 39.10	20 Ca 40.08	21 Sc 44.96	22 Ti 47.87	23 V 50.94	24 Cr 52.00	25 Mn 54.94	26 Fe 55.85	27 Co 58.93	28 Ni 58.69	29 Cu 63,55	30 Zn 65,38	31 Ga _{69.72}	32 Ge 72.64	33 As _{74.92}	34 Se _{78.96}	35 Br 79.90	36 Kr 83.80
37 Rb 85.47	38 Sr 87.62	39 Y 88.91	40 Zr 91.22	41 Nb 92.91	42 Mo 95.94	43 Tc	44 Ru 101.07	45 Rh	46 Pd 106.42	47 Ag 107.87	48 Cd 112.41	49 In	50 Sn	51 Sb 121.76	52 Te	53 126.90	54 Xe 131.29
55 Cs 132.91	56 Ba 137.33	57 La	72 Hf 178.49	73 Ta 180.95	74 W 183.84	75 Re	76 Os 190.23	77 r 192.22	78 Pt 195.08	79 Au 196.97	80 Hg _{200.59}	81 TI 204.38	82 Pb 207.20	83 Bi 208.98	84 Po (209)	85 At (210)	86 Rn (222)
87 Fr (223)	88 Ra (226)	89 Ac (227)	104 Rf (267)	105 Db (268)	106 Sg (269)	107 Bh (270)	108 Hs (270)	109 Mt (278)	110 Ds (281)	111 Rg (282)	112 Cn (285)	113 Nh (286)	114 FI (289)	115 Mc (290)	116 Lv (293)	117 Ts (294)	118 Og (294)

58	59	60	61	62	63	64	65	66	67	68	69	70	71
Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
140.12	140.91	144.24	(145)	150.36	151.96	157.25	158.93	162.50	164.93	167.26	168.93	173.04	174.97
90	91	92	93	94	95	96	97	98	99	100	101	102	103
Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr
232.04	231.04	238.03	(237)	(244)	(243)	(247)	(247)	(251)	(252)	(257)	(258)	(259)	(266)

232.04	231.04 238.03 (237) (244) (243)	(247) (247) (251) (252) (257) (258)	(259) (266)
constants	conversions	standard potentials at 25 °c	$C \qquad E^{\circ} (V)$
R = 0.08206 L atm/r	1 in = 2.54 cm	$F_2(g) + 2 e^- \rightleftharpoons 2 F^-(aq)$	+2.870 V
$R=8.314~\mathrm{J/mol~K}$	1 ft = 12 in	$Au^{+}(aq) + e^{-} \rightleftharpoons Au(s)$	+1.690 V
$F = 96485 \text{ C/mol e}^-$	1 yd = 3 ft	$Pd^{2+}(aq) + 2 e^{-} \rightleftharpoons Pd(s)$	+0.915 V
$N_{\rm A} = 6.022 \times 10^{23} / 10^{23}$	mol 1 mi = 5280 ft	$Ag^{+}(aq) + e^{-} \rightleftharpoons Ag(s)$	+0.800 V
$h = 6.626 \times 10^{-34} \text{ J} \cdot \text{s}$	1 lb = 453.6 g	$Fe^{3+}(aq) + e^{-} \rightleftharpoons Fe^{2+}(aq)$	+0.771 V
$c=3.00\times 10^8~\mathrm{m/s}$	1 ton = 2000 lbs	$Cu^{2+}(aq) + 2 e^{-} \rightleftharpoons Cu(s)$	+0.340 V
$g=9.81~\mathrm{m/s^2}$	1 tonne = 1000 kg	$2~\mathrm{H^+(aq)} + 2~\mathrm{e^-} ightleftharpoons H_2(\mathrm{g})$	0.000 V
	1 gal = 3.785 L	$Fe^{3+}(aq) + 3 e^{-} \rightleftharpoons Fe(s)$	-0.040 V
	$1 \text{ gal} = 231 \text{ in}^3$	$Pb^{2+}(aq) + 2 e^{-} \rightleftharpoons Pb(s)$	-0.130 V
conversions	1 gal = 128 fl oz	$Ni^{2+}(aq) + 2 e^{-} \rightleftharpoons Ni(s)$	-0.230 V
1 atm = 760 torr	1 fl oz = 29.57 mL	$Cr^{3+}(aq) + e^- \rightleftharpoons Cr^{2+}(aq)$	-0.407 V
1 atm = 101325 Pa	1.1.1.1.0	$Fe^{2+}(aq) + 2 e^{-} \rightleftharpoons Fe(s)$	-0.440 V
1 atm = 1.01325 bar		$Zn^{2+}(aq) + 2 e^{-} \rightleftharpoons Zn(s)$	-0.760 V
$1 \text{ bar} = 10^5 \text{ Pa}$	1 W = 1 J/s	$Cr^{2+}(aq) + 2 e^{-} \rightleftharpoons Cr(s)$	-0.910 V
$^{\circ}F = ^{\circ}C(1.8) + 32$	1 A = 1 C/s	$Mn^{2+}(aq) + 2 e^{-} \rightleftharpoons Mn(s)$	-1.180 V
$K = {^{\circ}C} + 273.15$		$Al^{3+}(aq) + 3 e^{-} \rightleftharpoons Al(s)$	-1.660 V
		$Mg^{2+}(aq) + 2 e^{-} \rightleftharpoons Mg(s)$	-2.360 V
		- (-)	

This exam should have exactly 25 questions. Each question is equally weighted at 4 points each. You will enter your answer choices on the virtual bubblehseet after you have finished. Your score is based on what you submit on the virtual bubblesheet and not what is circled on the exam.

- 1. What is the reducing agent in an alkaline cell?
- a. MnO_2 (s)
- b. Mn_2O_3 (s)
- c. ZnO (s)
- •d. Zn (s)
 - e. H₂O (ℓ)

Explanation: The overall reaction for an alkaline cell is $2MnO_2 + Zn \longrightarrow Mn_2O_3 + ZnO$ Zn changes its oxidation state from 0 to +2 and is thus oxidized and the reducing agent is the reactant that itself undergoes oxidation. It's the reactant that "gives away" electrons.

2. Balance the following redox reaction in acidic solution:

$$Ru^{3+} + IO_4^- \longrightarrow RuO_4 + I^-$$

What is the coefficient of Ru³⁺ in the simplest, overall reaction?

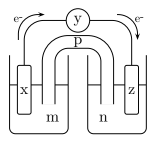
- a. 5
- b. 4
- ●c. 8
- d. 1
- e. 2
- f. 6

Explanation:

$$12\mathrm{H}_2\mathrm{O} + 8\mathrm{Ru}^{3+} + 5\mathrm{IO}_4^- \longrightarrow 8\mathrm{RuO}_4 + 5\mathrm{I}^- + 24\mathrm{H}^+$$

Coefficient on Ru^{3+} is 8.

3. (part 1 of 2): Consider the electrochemical cell diagram shown below which is running with the electron flow direction as shown. What must be true for this electrochemical cell to run a complete closed-loop (circular) circuit without creating an imbalance of charge?



- a. anions must flow from y toward z, cations flow from y toward x
- b. anions must flow from y toward x, cations flow from y toward z
- •c. anions must flow from p toward x, cations flow from p toward z
- d. anions must flow from p toward z, cations flow from p toward x
- e. anions must flow from p toward x, cations flow from y toward z

Explanation: p = salt bridge; cations will flow towards the cathode which has to be the z component due to the fact electrons are flowing towards it in the external circuit. Anions will flow to x, which is the anode compartment.

- **4.** (part 2 of 2): Suppose the previously shown cell diagram represents an electrolytic cell. What is the role and assigned charge of the component labeled *z*?
- a. anode, negative
- b. cathode, no charge
- c. cathode, positive
- •d. cathode, negative
 - e. anode, positive

Explanation: Electrons always travel from anode to cathode. For the diagram given, the electrons travel from x (anode) to z (cathode). For an electrolytic cell, the anode is positive (x) and the cathode is negative (z).

5. For the following reaction of titanium(III) chloride with fluorine, what is the oxidizing agent?

$$3\text{TiCl}_3(aq) + 3\text{F}_2(g) \rightarrow 3\text{TiF}_3(aq) + 3\text{Cl}_2(g)$$

- a. $Cl_2(g)$
- b. $TiF_3(aq)$
- c. TiCl₃(aq)
- d. Ti(aq)
- ●e. F₂(g)

Explanation: Fluorine is being reduced from an oxidation state of 0 to a state of -1, so it is causing the chloride ion to be oxidized.

6. Consider the cell:

$$Pt(s) \mid H_2(g) \mid H^+(aq) \mid \mid Cl^-(aq) \mid Cl_2(g) \mid C(s)$$

What is serving as the anode in this electrochemical cell?

- a. C(s)
- b. $Cl_2(g)$
- c. $H_2(g)$
- •d. Pt(s)
- $e. H^+(aq)$
- f. $Cl^{-}(aq)$

Explanation: When using standard cell notation, the solid to the furthest left side of the notation is solid substance serving as the anode.

7. What is the cell potential for the following nonstandard cell?

$$Ni \mid Ni^{2+} (1M) \mid | Ag^{+} (0.0065M) \mid Ag$$

- •a. 0.901 V
- b. 0.332 V
- c. 1.160 V
- d. 0.771 V
- e. 0.460 V
- f. 0.622 V

Explanation:
$$E^{\circ} = +0.800 - -0.230 = +1.03 \text{ V}.$$

 $Q = \frac{1}{0.0065^2}$ and n = 2

 $E = 1.03 - 0.05916/2\log(\frac{1}{0.0065^2})$

E = 0.901 V

- 8. A proposed solution for grid-scale energy storage is the use of so-called 'flow batteries'. They are essentially just warehouse sized electrochemical cells! They can be charged by running the cell electrolytically, and can later be discharged by running the cell voltaically. A common cell design uses the Cr^{2+}/Cr^{3+} and Fe^{2+}/Fe^{3+} redox couples. What is the standard cell potential for a voltaic cell constructed using these redox couples?
- a. +0.36 V
- b. 1.68 V
- ●c. +1.18 V
- d. -0.36 V
- e. -1.18 V
- f. 0.033 V

Explanation: The standard reduction potentials are -0.407 V for $\text{Cr}^{2+}/\text{Cr}^{3+}$ and $0.771 \text{ for Fe}^{2+}/\text{Fe}^{3+}$. We are running a voltaic cell, so we assign the more negative potential to be the anode $(\text{Cr}^{2+}/\text{Cr}^{3+})$ and the more positive potential to be the cathode $(\text{Fe}^{2+}/\text{Fe}^{3+})$, giving us a potential of +1.18 V.

- **9.** Car batteries from the 1970's were able to produce about 300 cold cranking amps. Modern car batteries are able to produce well over 600 cold cranking amps. This improvement in performance is mostly due to:
- a. The use of higher purity lead in batteries.
- b. Lead from the 1970's was just lazy.
- c. Batteries from the 1970's used PbO₂ as the oxidant, modern batteries now use TiO₂.
- d. Batteries from the 1970's used lead (II) sulfate, modern batteries now use lead (II) nitrate.
- e. The design of improved electrodes with higher surface area.

Explanation: Car batteries from the 1970's and modern car batteries operate off of the exact same electrochemical reaction. The improvement in current output is largely due to the design improved electrodes with increased surface area.

- 10. Which of the following species is the strongest reducing agent?
- •a. Zn(s)
 - b. Ag(s)
 - c. Cu(s)
 - d. $Mg^{2+}(aq)$
 - e. $F_2(g)$

Explanation: A good reducing agent needs to push electrons away which means they will be on the right side of a standard potential table and will have the most negative E° . That means that Zn is the best of those listed.

- 11. I put a piece of copper wire into a solution of 1 M Ag⁺. What happens?
- a. The two reactants will fuse together to make Zn metal.
- b. Nothing will happen.
- c. The copper ions in solution are reduced and plate onto the copper wire
- •d. The copper wire reduces the Ag⁺ ions to silver metal.

Explanation: The standard potential for Cu(s) and Ag⁺(aq) is 0.46 V. This is spontaneous, so the copper wire will be oxidized and the silver ions will be reduced.

- 12. You hook up a voltmeter to a standard electrochemical cell and it reads +0.313 V. With this information, which of the following is true regarding this cell?
- a. ΔG is positive and K is less than 1.
- •b. ΔG is negative and K is greater than 1.
 - c. ΔG is positive and K is greater than 1.
 - d. ΔG is negative and K is less than 1.
 - e. ΔG is zero and K is equal to 1.

Explanation: Recall the triforce, $-nF\mathcal{E}^{\circ} = \Delta G^{\circ} = -RTlnK$. If we measure a positive potential, this must correspond to a negative ΔG° (a spontaneous process). This must also correspond to an equilibrium constant which is greater than 1.

- 13. Salts of chromate $(\operatorname{CrO}_4^{2-})$ are incredibly toxic and carcinogenic because they mimic phosphate $(\operatorname{PO}_4^{3-})$, which forms backbone of DNA. What is the oxidation number of Cr in chromate and P in phosphate, respectively?
- a. +8 and +8
- •b. +6 and +5
- c. +5 and +5
- d. +5 and +6
- e. +6 and +6

Explanation: For CrO_4^{2-} , each oxygen is 2-, but the overall complex has a 2- charge, making the Cr 6+. For PO_4^{3-} , each oxygen is again -2, but this time the overall molecule has a 3- charge, making the P 5+.

14. We construct the following cell below and measure the potential to be 1.610 V. What is the concentration of $\mathrm{Mn^{2+}}$ ions?

$$Mn \mid Mn^{2+} (? M) \mid | Cu^{2+}(2M) \mid Cu$$

- a. 0.256 M
- •b. 0.0018 M
 - c. 0.767 M
 - d. 0.110 M
 - e. 0.0073 M

Explanation: Use Nernst equation:

$$\begin{split} E &= E^{\circ} - \frac{0.05916}{2} \log Q \\ E^{\circ} &= E^{\circ}_{cathode} - E^{\circ}_{anode} = 0.340V - (-1.180V) = 1.520V \\ 1.610 &= 1.520 - \frac{0.05916}{2} \log(\frac{x}{2}) \\ \text{Solve for } x = 0.0020 \text{ M} \end{split}$$

- 15. Which of the following metals is most easily oxidized?
- a. Ag
- b. Cu
- c. Au
- d. Ni
- ●e. Al

Explanation: The metal that is most easily oxidized is the one with the greatest oxidation potential. The table provided shows reduction potentials for ions associated with these metals. The oxidation potential of the metal is the opposite of the standard reduction potential of the associated ions.

- 16. In an electrolytic cell, the positive terminal is the _____ and is the site of the _____ half-reaction.
- •a. anode; oxidation
 - b. anode; reduction
 - c. cathode; oxidation
 - d. cathode; reduction

Explanation: In an electrolytic cell (analogous to charging a battery), the cathode is always attributed a negative sign and the anode a positive sign. By definition, the cathode is the site of reduction, and the anode is the site of oxidation.

17. Consider the following redox reaction involving curium (An element whose namesake honors Marie Curie, a pioneering chemist and physicist who studied radioactivity) and the unfortunately named Uranate ion, $\mathrm{UO}_4^{2^-}$.

$$Cm(s) + UO_4^{2-}(aq) \longrightarrow Cm^{3+}(aq) + UO_2(s)$$

Using the smallest possible integer coefficients, balance the reaction in acidic aqueous solution. What is the coefficient for H⁺, and which side of the reaction is it on?

- a. 10, left
- b. 2, left
- c. 12, right
- d. 10, right
- e. 4, left
- •f. 12, left

Explanation:

$$2\mathrm{Cm} + 3\mathrm{UO_4^{2-}} + 12\mathrm{H^+} \longrightarrow 2\mathrm{Cm^{3+}} + 3\mathrm{UO_2} + 6\mathrm{H_2O}$$

Coefficient on H⁺ is 12, and is on the left.

- 18. Which type of battery chemistry listed below results in a battery that is not rechargeable?
- a. Li-ion
- b. NiMH

explin: NiCd, Pb-acid, Li-ion, and NiMH are all examples of secondary cells (rechargeable) batteries. The only one that is a primary (non-rechargable) battery is an alkaline cell.

- c. Pb-acid
- d. alkaline
- e. NiCd

Explanation:

19. Consider the following redox reaction:

$$Ta^{4+}(aq) + Co(s) \longrightarrow Ta^{2+}(aq) + Co^{2+}(aq)$$

What is the reducing agent?

- •a. Co
 - b. Co^{2+}
 - c. Ta^{2+}
 - d. Ta⁴⁺

Explanation: The reducing agent is the species being oxidized, which is Co(s) being oxidized to Co^{2+} .

20. You go to the hardware store to buy some root killer (copper(II) sulfate, 159.6 g/mol) and to your dismay it is all sold out. Using your electrochemistry knowledge, you decide to make some yourself. You instead buy copper piping and sulfuric acid to construct an electrolytic cell. Copper(II) sulfate is deposited on the copper electrode from a solution of sulfuric acid. What mass of copper(II) sulfate is made when 27.0 amps of current is run continuously for 17.0 hours?

- a. 22.8 g
- b. 672 g
- ●c. 1.37 kg
 - d. 308 g
- e. 1.76 kg
- f. 1.13 kg

Explanation: The oxidation number of copper sulfate is +2, so you need to lose 2 mols of electrons per mol of copper to make Cu(II). Now use $mol = \frac{I \cdot t}{n \cdot F} = \frac{(27) \cdot (17*60*60)}{2.96458} = 8.56$ mols of CuSO₄. Use MW to convert into g and then into kg to find 1.37 kg.

- 21. Which of the following is true of a standard hydrogen electrode?
 - I. It cannot be part of a voltaic cell since its standard reduction potential is zero.
 - II. It contains a solution with a concentration of $[\mathrm{H^+}] = 1~\mathrm{M}$
- III. It uses H_2 gas at a pressure of 1 atm.
- a. I and II
- b. I and III
- c. only I
- •d. II and III
 - e. only III
 - f. I, II and III
 - g. only II

Explanation: SHE is used as a reference for all other half-reactions, and is zero by definition. That does not mean that it cannot be used in a cell. By definition, the solution must have 1 M $\rm H^+$ and the $\rm H_2$ pressure must be 1 atm. Thus, the correct answers are only II and III.

22. What is the standard free energy change for the following standard cell?

Pb | Pb
$$^{2+}$$
 (1M) || Cu $^{2+}$ (1M) | Cu

- a. +45.3 kJ
- b. +90.7 kJ
- c. +40.5 kJ
- d. -45.3 kJ
- e. −90.7 kJ
- f. -40.5 kJ

Explanation: We can relate the cell potential to the standard free energy change using one of the triforce relationships: $-nF\mathcal{E}^{\circ} = \Delta G^{\circ}$.

Recognize that Pb is the anode and Cu is the cathode:

$$\mathcal{E}^\circ = \mathcal{E}_{\text{constant}}^\circ - \mathcal{E}_{\text{constant}}^\circ - \mathcal{E}_{\text{constant}}^\circ = 0.340 \ V$$
 - (-0.130 V) = 0.470 V

Rearrange the above realtionship and solve for $\Delta G^{\circ} =$ -90.7 kJ

- 23. How long will it take to deposit 0.0231 moles of solid promethium (Pm, atomic number 61) by the electrolysis of $Pm(NO_3)_3(aq)$ using a current of 1.31 A?
- a. 70 minutes
- b. 60 minutes
- •c. 85 minutes
 - d. 75 minutes
 - e. 80 minutes
- f. 65 minutes

Explanation: First note that the oxidation number of promethium is +3 in the aqueous solution given, so when it gets reduced to solid promethium, 3 moles of electrons will be transferred. Now use $t = \frac{mol \cdot n \cdot F}{I} = \frac{0.0231(3)(96485)}{1.31} = 5100 \text{ s} = 85 \text{ minutes}$

24. The following concentration cell has a measured potential of +84.8 mV. What is the unknown chromium(II) ion concentration in this cell?

$$Cr \mid Cr^{2+} (1.7 \times 10^{-4} M) \mid | Cr^{2+} (? M) \mid Cr$$

- a. $3.2 \times 10^{-8} \text{ M}$
- b. 0.14 M
- c. 0.0023 M
- d. 0.075 M
- e. $2.0 \times 10^{-6} \text{ M}$
- •f. 0.125 M

Explanation: Use Nernst equation:

$$E = E^{\circ} - \frac{0.05916}{2} \log Q$$

$$0.0848 = 0 - (0.02958) \log(\frac{1.7 \times 10^{-4}}{x})$$

solve for x = 0.125 M = $[Cr^{2+}]$ in the cathode compartment

- 25. Identify the oxidation number for each element in MnSO₄. List your answer in the order Mn, S, and O.
- a. +2, +5, -2
- b. +2, +8, -2
- c. +4, +3, -1
- •d. +2, +6, -2
 - e. +2, +4, -6

Explanation: Oxygen has an oxidation number of -2. The polyatomic sulfate anion has an overall charge of -2, so sulfurs's oxidation number be -6 in order to compensate for the four oxygen ions. The manganese is given as manganese(II) ion, so its oxidation number in considered to be +2.

After you are finished and have all your answers circled, go to the front of the room and then use the QR code show below to pull up the virtual answer page for your exam. Enter the appropriate info plus all your answers - click the SUBMIT button. Double check your choices on the next page. Once your are sure, click the submit button on that page to enter your answers. Make sure you get the confirmation screen (different background color!) and show it to the TA or proctor. After that, turn in your exam and scratch paper. You're free to leave after that.



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