

#### Objective Questions

#### **Electrolytes and Electrolysis**

1.	Which	of	the	following	will	not	conduct	electricity	in	aqueous
	solution	1						[/	Mu	1982, 83]

- (a) Copper sulphate
- (b) Sugar
- (c) Common salt
- None of these (d)
- 2. Strong electrolytes are those which

[MNR 1983]

- (a) Dissolve readily in water
- Conduct electricity
- (c) Dissociate into ions at high dilution
- (d) Completely dissociate into ions at all dilutions
- In aqueous solution, strong electrolytes [AMU 1983, 84] 3.
  - (a) Are partially ionized
  - (b) Do not ionise
  - Ionise almost completely (c)
  - (d) Form polymers

An electrolyte

[KCET 1984; MP PET/PMT 1988]

- (a) Forms complex ions in solution
- Gives ions only when electricity is passed
- Possesses ions even in solid state (c)
- 5.
  - (a) They are unstable
  - (b) The water dissolves it
  - The force of repulsion increases
  - (d) The forces of electrostatic attraction are broken down by water
- Electrolyte can conduct electricity because
  - (a) Their molecules contain unpaired electrons, which are mobile
  - (b) Their molecules contain loosely held electrons which get free under the influence of voltage
  - The molecules break up into ions when a voltage is applied
  - The molecules are broken up into ions when the electrolyte is fused or is dissolved in the solvent
- one of the following metals could not be obtained on 7. electrolysis of aqueous solution of its salts [IIT 1990]
  - (a) Ag
- (b) Mg
- (c) Cu
- (d) *Cr*
- Which of the following aqueous solution will conduct an electric current quite well [MP PMT 1987]
  - Glycerol
- (b) HCl
- Sugar
- (d) Pure water
- On the electrolysis of aqueous solution of sodium sulphate, on 9. [MP PMT 1992, 2002] cathode we get
  - (a) Na
- (b)  $H_2$
- (c)  $SO_2$
- (d)  $SO_3$
- 10. Electrolysis involves oxidation and reduction respectively at

#### [CPMT 1973; AMU 1983; NCERT 1983, 84; MH CET 2001]

- (a) Anode and cathode
- (b) Cathode and anode
- (c) At both the electrodes
- (d) None of the above
- Which of the following compounds will not undergo decomposition 11. on passing electricity through aqueous solution
  - (a) Sugar
- (b) Sodium Chloride
- (c) Sodium Bromide
- (d) Sodium Acetate
- 12. During the electrolysis of an electrolyte, the number of ions produced, is directly proportional to the [AFMC 2002]

- Time consumed
- Electro chemical equivalent of electrolysis (b)
- (c) Quantity of electricity passed
- (d) Mass of electrons
- When the sample of copper with zinc impurity is to be purified by 13. electrolysis, the appropriate electrodes are

[AIEEE 2002]

Cathode Anode Pure copper (a) Pure zinc (b) Impure sample Pure copper Impure zinc Impure sample (d) Pure copper Impure sample

In the electrolytic cell, flow of electrons is from

[IIT Screening 2003]

- (a) Cathode to anode in solution
- (b) Cathode to anode through external supply
- Cathode to anode through internal supply
- Anode to cathode through internal supply
- An electric current is passed through an aqueous solution of the 15. following. Which one shall decompose [NCERT 1972]
  - (a) Urea
- (b) Glucose
- (c)  $AgNO_3$
- (d) Ethyl alcohol
- 16. The electric conduction of a salt solution in water depends on the
  - Shape of its molecules
  - Size of its molecules (b)
  - Size of solvent molecules (c)

(d) Gives ions only when dissolved in water

(d) Extent of its ionization

Electrolytes when dissolved in water dissociates into ions because [CPMT 1974, 78; MNR 1983]

A solution of sodium sulphate in water is electrolysed using inert electrodes. The products at the cathode and anode are respectively[IIT 1987,96]

- (a)  $H_2$ ,  $O_2$
- (b)  $O_2, H_2$
- (c)  $O_2$ , Na
- (d)  $O_2$ ,  $SO_2$
- On electrolysing a solution of dilute  $H_2SO_4$  between platinum 18. electrodes, the gas evolved at the anode is

[NCERT 1977, 79; MNR 1980; CBSE PMT 1992]

- (a)  $SO_2$
- (b)  $SO_2$
- (c)  $O_2$
- (d)  $H_2$
- The addition of a polar solvent to a solid electrolyte results in 19.
  - (a) Polarization
- (b) Association
- (c) lonization
- (d) Non-liberation of heat
- 20. During the electrolysis of fused NaCl, which reaction occurs at anode [NCERT 1973; AFMC 1992; MP PMT 2002]
  - (a) Chloride ions are oxidized
  - Chloride ions are reduced
  - Sodium ions are oxidised
  - Sodium ions are reduced
- 21. The amount of ion discharged during electrolysis is not directly proportional to [NCERT 1973]
  - (a) Resistance
  - (b) Time
  - Current
  - Chemical equivalent of the ion
- Electrolysis of aqueous HCl solution produces 22.

[CPMT 1987]

- HMPgPETIt200e anode
- $H_2$  gas at the cathode
- $Cl_2$  gas at the cathode



[EAMCET 1990; MP PET 1994, 97]

(b) 107.87 g

Hydrogen is liberated at the cathode (d)  $Cl_2$  and  $O_2$  gases both at the anode (b) Hydrogen is liberated at the anode During electrolysis of NaCl solution, part of the reaction is 23. There is no reaction  $Na^+ + e^- \rightarrow Na$ . This is termed as [NCERT 1984]  $H^-$  ions produced migrate to the cathode (a) Oxidation (b) Reduction During electrolysis, the species discharged at cathode are 35. (c) Deposition (d) Cathode reaction [AFMC 2000] When a solution of an electrolyte is heated the conductance of the 24. (b) Cation (a) lons [KCET 1991] (d) All of these (c) Anion (a) Increases because of the electrolyte conducts better (b) Decreases because of the increased heat 36. Electrolysis of molten anhydrous calcium chloride produces Decreases because of the dissociation of the electrolyte is [AIIMS 2000] suppressed (a) Calcium (b) Phosphorus (d) Increases because the electrolyte is dissociated more (c) Sulphur (d) Sodium The passage of current liberates  $H_2$  at cathode and  $Cl_2$  at anode. 25. 37. Which of the following properties of pure metal makes it more useful then the corresponding alloy [RPET 2000] The solution is [EAMCET 1979,87] (a) It is harder than corresponding alloy (a) Copper chloride in water (b) NaCl in water (b) It has high density  $H_2SO_4$ (d) Water (c) It can be extracted easily 26. Pure water does not conduct electricity because it (d) It conducts heat and electricity easily [Manipal MEE 1995] 38. Which of the following liberate hydrogen on reaction with dilute (a) Has a low boiling point  $H_2SO_4$ [Roorkee 2000] (b) Is almost totally unionized (a) *Fe* (c) Is neutral Си (d) Is readily decomposed (c) Al (d) Hg27. Which is responsible for electrical conduction of molten sodium Which one of the following material conducts electricity 39. chloride [MADT Bihar 1995] [Kerala (Med.) 2003] (a) Free electrons (b) Free ions Crystalline sodium chloride (c) Free molecules Barium sulphate (d) Atoms of sodium and chlorine Fused potassium chloride In electrolysis of aqueous copper sulphate, the gas at anode and 28 cathode is Molten sulphur (b)  $SO_2$  and  $H_2$ (a)  $O_2$  and  $H_2$ Which of the following metals will give  $\,H_{2}\,$  on reaction with NaOH 40. (d)  $SO_3$  and  $O_2$ (c)  $H_2$  and  $O_2$ (a) Mg (b) *Ba* Use of electrolysis is (c) Ca (d) Sr [AFMC 1995] 29. Which of the following is not a non electrolyte [] & K 2005] (a) Electroplating (b) Electrorefining (a) Acetic acid (b) Glucose (a) and (b) both (d) None of these (c) Ethanol (d) Urea Sodium is made by the electrolysis of a molten mixture of about 30. 40% NaCl and 60% CaCl2 because Faraday's law of electrolysis [CBSE PMT 1995] CaCl<sub>2</sub> helps in conduction of electricity Amount of electricity that can deposit 108 gm of silver from AgNO 1. (b) This mixture has a lower melting point than NaCl solution is [AFMC 1993; MP PMT 2004] (a) 1 ampere (b) 1 coulomb Ca<sup>++</sup> can displace Na from NaCl (c) 1 faraday (d) None of the above (d) Ca++ can reduce NaCl to Na When 9.65 coulombs of electricity is passed through a solution of Electrolysis is a process in which the cations and anions of the 31. silver nitrate (atomic weight of Ag = 107.87 taking as 108) the electrolyte are [MP PET 1995] amount of silver deposited is (b) Hydrolysed (a) Hydrated [EAMCET 1992; KCET 2000] (c) Charged (d) Discharged (a) 10.8 mg (b) 5.4 mg Degree of ionisation of a solution depends upon 32. (c) 16.2 mg (d) 21.2 mg [BHU 1998] Three faradays electricity was passed through an aqueous solution of iron (II) bromide. The weight of iron metal (at. wt. = 56) deposited (b) Nature of the electrolyte (a) Temperature at the cathode (in gm) is (c) Nature of the solvent (d) None of these [EAMCET 1991] Which of the following is non-electrolytes 33. (a) 56 (b) 84 [KCET (Med.) 1999] (c) 112 (d) 168 NaCl (b)  $CaCl_2$ (a) A silver cup is plated with silver by passing 965 coulombs of electricity, the amount of silver deposited is

[JIPMER 1999]

(a) 9.89 g

(d) CH<sub>3</sub>COOH

(c)  $C_{12}H_{22}O_{11}$ 

34.

When a molten ionic hydride is electrolysed



16.

17.

(a) 20.5

 $Al^{3+}$ 

(a)  $3 \times 96500 \ C$ 

Co   10787 g	UNIVE		chemistry								
passed through a solution of $AI^{***}$ ions, the weight of $AI$ deposited is $(NCRRT 1984]$ (d) $36 \ gm$ (b) $36 \ gm$ (c) $45 \ gm$ (d) $39 \ gm$ (h) $36 \ gm$ (d) $39 \ gm$ (a) $39 \ gm$ (d) $39 \ gm$ (d) $39 \ gm$ (d) $39 \ gm$ (d) $39 \ gm$ (e) Coulometer (e) Coulometer (d) Colorimeter (E) Coulometer (d) Colorimeter (E) Coulometer (e) Gram/coulomb (d) Coulomb/gram (e) Gram/coulomb (e) Gram/coulomb (d) Coulomb/gram (e) Gram/coulomb (e) Gram/coulomb (d) Coulomb/gram (e) Gram/coulomb (e) Gram/coulomb (e) Gram/coulomb (f) Gram/coulomb (g) Gr		(c) 1.0787 g	(d) 1.002 g				96500 ~			96500 ~	
deposited is $(3) \ 27 \ gm$ (b) 36 $gm$ (b) 36 $gm$ (c) 45 $gm$ (d) 39 $gm$ (e) 45 $gm$ (d) 39 $gm$ (e) 45 $gm$ (d) 39 $gm$ (a) Calorimeter (b) Cathetometer (c) Coulometer (d) Colorimeter (d) Colorimeter (e) Coulometer (d) Colorimeter (e) Gamicoulomb (d) Coulombigeam (e) Colorimeter (e		The atomic weight of	Al is 27. When a curren	t of 5 Faradays is		(c)	$\frac{}{3}$ $C$		(d)	$\frac{}{2}$	
deposited is $(3) \ 27 \ gm$ (b) 36 $gm$ (b) 36 $gm$ (c) 45 $gm$ (d) 39 $gm$ (e) 45 $gm$ (d) 39 $gm$ (e) 45 $gm$ (d) 39 $gm$ (a) Calorimeter (b) Cathetometer (c) Coulometer (d) Colorimeter (d) Colorimeter (e) Coulometer (d) Colorimeter (e) Gamicoulomb (d) Coulombigeam (e) Colorimeter (e		passed through a so	lution of $Al^{+++}$ ions, the	e weight of <i>Al</i>	18.	On 1	passing one fa	raday of elec	tricity	through the	electrolytic cells
(a) $27 \ gm$ (b) $36 \ gm$ (d) $39 \ gm$ (e) Coulometer (d) Coulometer (d) Colometer				-				•	•	•	•
(c) 43 gm		(a) 27 gm	(b) 36 <i>gm</i>				•				•
(a) Calorimeter (b) Cathetometer (c) Coulometer (d) Colorimeter (d) Colorimeter (d) Colorimeter (d) Colorimeter (e) Coulomb (e) Gram/coulomb (d) Combination (d) Combination (e) Gram/coulomb (d) Combination		(c) 45 gm	(d) 39 gm			Ag(	(At. wt. = 10a)	5), Ni (At.wi	$x_{\cdot} = 5$	9) and $Cr(A)$	t.Wt. = 32) is[AllM
(a) Claylimeter (b) Cathetometer (c) Coulomber (d) Colorimeter (e) Colorimeter (d) Colorimeter (EMMCET 1980) [RAMCET 1980] (a) $Gram$ (b) $Gram/ampere$ (c) $Gram/ampere$ (d) $Gram/ampere$ (e) $Gram/ampere$ (f) $Gram/ampere$ (g) $Gram/ampere$ (h) $Gram/ampere$ (b) $Gram/ampere$ (b) $Gram/ampere$ (c) $Gram/ampere$ (d) $Gram/ampere$ (e) $Gram/ampere$ (f) $Gram/ampere$ (h)			the measurement of quant			(2)					
(c) Coulometer  (d) Colorimeter  The unit of electrochemical equivalent is  [EAMCET 1980]  (a) Gram (b) Gram/ampere (c) Gram/coulomb (d) Coulomb/gram  (d) Goal gam (b) Gram/ampere (e) Gram/coulomb (d) Coulomb/gram  A certain current liberated to 30.98 gam (b) 105.09 gm (c) 20.18 gm (d) 63.5 gm (d) 127.0 gm (e) 62.5 gm (d) 127.0 gm (e) 63.5 gm (d) 127.0 gm (e) 63.5 gm (d) 127.0 gm (e) 63.5 gm (d) 127.0 gm (e) Anions move towards anode, cations towards cathode (b) Anions move towards cathode, cations towards anode (d) No movement of ions takes place  Unit of Faraday is  (c) Coulomb $mole^{-1}$ (d) Coulomb $Sec^{-1}$ On passing 0.1 Faraday of electricity through aluminium chloride, the amount of aluminium metal deposited on cathode is $(Al = 27)$ [MP PMT 1991]  (a) $0.9 gm$ (b) $0.3 gm$ (c) $0.3 gm$ (d) $0.3 gm$ (e) $0.3 gm$ (b) $0.3 gm$ (c) $0.27 gm$ (d) $0.27 gm$ (d) $0.27 gm$ (e) $0.3 gm$ (b) $0.3 gm$ (c) $0.4065 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (d) $0.50.0 gm$ (e) $0.4065 gm$ (f) $0.50.0 gm$ (f) $0.50.0 gm$ (f) $0.50.0 gm$ (g) $0.50.0 gm$ (g		(a) Calorimeter	(b) Cathetomet	ter		. ,	•	_		_	
The unit of electrochemical equivalent is [EAMCET 1980] (a) $Gram$ (b) $Gram/ampre$ (c) $Gram/coulomb$ (d) $Goulombsyram$ (d) $Goulombsyram$ (a) $Goulombsyram$ (a) $Goulombsyram$ (b) $Goulombsyram$ (b) $Goulombsyram$ (b) $Goulombsyram$ (b) $Goulombsyram$ (c) $Goulombsyram$ (d) $Goulombsyram$ (e) $Goulombsyram$ (d) $Goulombsyram$ (e) $Goulombsyram$ (d) $Goulombsyram$ (e) $Goulombsyram$ (d) $Goulombsyram$ (e)		(c) Coulometer	(d) Colorimeter	r				_		-	
(a) Gram (b) Gram/ampere (c) Gram/coulomb (d) Coulomb/gram A certain current liberated 0,504 gm of hydrogen in 2 hours. How many grams of copper can be liberated by the same current flowing for the same time in a copper sulphate solution (NECRT 1973, 77; CPMT 1979, 88, All/MS 1954 points weight of copper will be deposited by passing 2 Faradays of electricity through a cupric salt (Atomic weight of Cu = 63.5) [NCERT 1975; CPMT 1979, 88, All/MS 1954 points weight (a) 20 gm (b) 3,75 gm (d) 63.5 gm (d) 12.70 gm  If the current is passed into the solution of an electrolyte (a) Anions move towards anode, cations towards anode (b) Anions and cations both move towards anode (c) Anions move towards atode, cations towards anode (d) No movement of ions takes place Unit of faraday is (c) Coulomb mole (d) Coulomb Sec (o) (n) Amper (b) Coulomb (c) Coulomb mole (d) Coulomb Sec (e) (a) 2.7 gm (b) 0.3 gm (c) 2.7 gm (d) 2.7 gm (d) 2.7 gm (d) 2.7 gm (d) 2.7 gm (e) 6.8 gm (e) 0.4065 gm (d) 40.55 gm (d) 40.65 gm (e) 0.4065 gm (d) 40.65 gm (d) 40.65 gm (d) 40.65 gm (e) 0.4065 gm (e) 0.4065 gm (e) 0.4065 gm (f) 40.65 gm (g) 40.6		The unit of electrocher	nical equivalent is	[EAMCET 1980]							
So the same deposits of the same current flowing for the same time in a copper sulphate solution (NCERT 1973, 77; CPMT 1979, 89; AllWei1994prolecule of Cu (d) 1 $gm$ quivalent of $Cu$ when 1 coulomb of charge is passed through electrolyte solution, then the mass deposited is equal to (a) $2.0 \ gm$ (b) $3.5 \ gm$ (d) $63.5 \ gm$ (d) $127.0 \ gm$ (b) $137.5 \ gm$ (d) $127.0 \ gm$ (d) $127.0 \ gm$ (e) $33.5 \ gm$ (d) $127.0 \ gm$ (h) $137.5 \ gm$ (e) $33.5 \ gm$ (d) $127.0 \ gm$ (h) $127.0 \ gm$ (g) $127.0 \ gm$ (h) $127.0 \ gm$ (h) $127.0 \ gm$ (g) $127.0 \ gm$ (h) $127.0 \ gm$ (h) $127.0 \ gm$ (h) $127.0 \ gm$ (h) $127.0 \ gm$ (l) $127.0 \ gm$ (a) $127.0 \ gm$ (b) $127.0 \ gm$ (a) $127.0 \ gm$ (b) $127.0 \ gm$ (b) $127.0 \ gm$ (a) Anions move towards anode, cations towards anode (d) No movement of ions takes place (c) Anions move towards achode, cations towards anode (d) No movement of ions takes place (e) Coulomb $127.0 \ gm$ (b) $127.0 \ gm$ (c) $127.0 \ gm$ (e) $127.0 \ gm$ (f) $127.0 \ gm$ (h) $127.0 \ gm$		( ) -			10			_			.l:C
A certain current liberated 0.504 $gm$ of hydrogen in 2 hours. How many grams of copper can be liberated by the same current flowing for the same time in a copper sulphate solution (NCERT 1973, 77; CPMT 1979, 89; Alli&i 1989) policially a cupric salt (Atomic weight of Cu = 63.5) [NCERT 1975; CPMT 1977]. Pg/ornic weight (2) 2.0 $gm$ (b) 3.75 $gm$ (d) 12.70 $gm$ (b) 3.75 $gm$ (d) 12.70 $gm$ (l) 12.70 $gm$ (l) 13.75 $gm$ (AlliMS 1979) [AlliMS 1979] (a) Anions move towards anode, cations towards cathode (b) Anions move towards anode, cations towards anode (c) Anions move towards cathode, cations towards anode (d) No movement of ions takes place (l) Columb $gm$ (b) 0.3 $gm$ (b) 0.3 $gm$ (c) 0.40 $gm$ (b) 0.3 $gm$ (d) 0.5 $gm$ (e) 0.40 $gm$ (b) 0.3 $gm$ (b) 0.3 $gm$ (c) 0.405 $gm$ (d) $gm$ (b) $gm$ (d) 4.055 $gm$ (e) 0.4065 $gm$ (f) 4.065 $gm$ (g) 4.40 $gm$ (h) 4.72 $gm$ (h) 4.72 $gm$ (h) 5.70 $gm$ (c) 0.4065 $gm$ (d) 4.72 $gm$ (d) 4.72 $gm$ (d) 4.72 $gm$ (e) 0.4065 $gm$ (e) 0.4065 $gm$ (f) 0.40 $gm$ (f) 0.50 $gm$ (f) 0.50 $gm$ (g) 4.72 $gm$ (f) 6.50.4 $gm$ (g) 6.50.4 $gm$ (f) 6.50.4 $gm$ (g) 6.50.4 $gm$ (			•		19.			ctricity when	passe	a through a so	
many grams of copper can be liberated by the same current flowing for the same time in a copper sulphate solution [NCERT 1973, 77; CPMT 1978, 89; All\( \) 89\( \) 89\( \) 99\( \) 99\( \) 60\( \) 127\( \) gm \\ (d) 63.5\( \) gm \\ (d) 63.5\( \) gm \\ (d) 127.9\( \) gm \\ (d) 127.0\( \) gm \\ (d) 127.0\			, ,			•	•				
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What weight of copper will be deposited by passing 2 Faradays of electricity through a cupric salf (Atomic weight of $Cu$ = 63.5 [NCERT 1975; CPMT 1975], 76/mornic weight (2) and (3) 2.0 gm (b) 3.175 gm (c) 63.5 gm (d) 127.0 gm (Earth of the current is passed into the solution of an electrolyte [AIIMS 1979] (a) Annions move towards anode, cations towards cathode (b) Anions move towards cathode, cations towards anode (c) Anions move towards cathode, cations towards anode (d) No movement of ions takes place [Unit of Faraday is [MP PMT 1991] (a) $A = 100$ $A = 1$				2011 1973, 77, CI WI 1975					. ,		
Equivalent weight of copper will be deposited by passing 2 Faradays of electricity through a cupric salt (Atomic weight of $Cu = 63.5$ ) [NCERT 1975; CPMT 1977]. A formic weight of $Cu = 63.5$ ] (a) 2.0 gm (b) 3.775 gm (c) 63.5 gm (d) 127.0 gm (d) 127.0 gm (a) 127.0 gm (a) 127.0 gm (b) 127.0 gm (a) 127.0 gm (b) 127.0 gm (a) 127.0 gm (b) 127.0 gm (b) 127.0 gm (c) Anions move towards anode (c) Anions move towards anode (c) Anions move towards anode (d) No movement of ions takes place (c) Coulomb mote $a_0$ (d) Coulomb $a_0$ (e) Coulomb $a_0$ (e) Coulomb $a_0$ (e) Coulomb $a_0$ (e) O.27 gm (d) 2.7 gm (d) 2.7 gm (d) 2.7 gm (d) 2.7 gm (e) 0.27 gm (d) 2.7 gm (f) 2.7 gm			. , .					. •		a timough ele	cerolyte solution,
What weight of copper will be deposited by $Cu = 63.5$ [NCERT 1975; CPMT 1975]. Atomic weight (a) 2.0 gm (b) 3.175 gm (c) 63.5 gm (d) 127.0 gm (d) 127.0 gm (d) 127.0 gm (e) 13.175 gm (d) 127.0 gm (e) 13.175 gm (		.,		1 6			•	•			
(c) $63.5\ gm$ (b) $3.75\ gm$ (c) $63.5\ gm$ (d) $127.0\ gm$ If the current is passed into the solution of an electrolyte [AllMS 1979]  (a) Anions move towards anode, cations towards anode (b) Anions move towards anode, cations towards anode (c) Anions move towards cathode, cations towards anode (d) No movement of ions takes place  Unit of Faraday is  (c) Coulomb $mole^{-1}$ (d) Coulomb $Sec^{-1}$ On passing 0.1 Faraday of electricity through aluminium chloride, the amount of aluminium metal deposited on cathode is $(Al = 27)$ [MP PMT 1991]  (a) $0.9\ gm$ (b) $0.3\ gm$ (c) $0.2\ rg$ m $0.1\ 2.7\ gm$ Which of the following represents the first law of Faraday  Which of the following represents the first law of Faraday  (c) $m = ect$ (d) $PV = nRT$ 5 amperes is passed through a solution of zinc sulphate for 40 minutes. Find the amount of zinc deposited at the cathode (CBSE PMT 1996]  (a) $40.65\ gm$ (b) $4.065\ gm$ (c) $0.4065\ gm$ (d) $65.04\ gm$ In an electroplating experiment $m\ g\ of\ silver\ deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991]  (a) 4m (b) m/2  (b) m/2  (c) m (d) m (d) m (e) m (e) m (b) m/2  (a) m (b) m/2  (b) m (c) m (d) m (d) m (e) m (e) m (d) m (e) m (e) m (e) m (for m)) of silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991]$		What weight of coppe	r will be deposited by pass	sing 2 Faradays of	. CDMT						
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sulphate solution, the amount of electricity required (in coulombs) is sulphate solution, the amount of electricity required (in coulombs) is (a) 40.65 gm (b) 4.065 gm (d) 65.04 gm (e) 0.4065 gm (e) 0.4065 gm (d) 1930 (e) 10 electrolysis of a fused salt, the weight of the deposit on an electrode will not depend on (a) Temperature of the bath (b) Current intensity (c) Electrochemical equivalent of ions (d) Time for electrolysis		( )	( )		24	. ,		am of conn	` '	-	f aqueous cupric
(a) 40.65 gm (b) 4.065 gm (c) 0.4065 gm (d) 65.04 gm In an electroplating experiment m g of silver is deposited, when 4 amperes of current flows for 2 minutes. The amount (in gms) of silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991] (a) 4m (b) m/2  (c) 3860 (d) 1930  In electrolysis of a fused salt, the weight of the deposit on an electrode will not depend on (a) Temperature of the bath (b) Current intensity (c) Electrochemical equivalent of ions (d) Time for electrolysis			int of zinc deposited at the		-	sulpl	nate solution, t		elect	ricity required	
In an electroplating experiment $m$ $g$ of silver is deposited, when 4 amperes of current flows for 2 minutes. The amount (in $gms$ ) of silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991]  (a) $4m$ (b) $m/2$ 25. In electrolysis of a fused salt, the weight of the deposit on an electrode will not depend on [CPMT 1973]  (a) Temperature of the bath  (b) Current intensity  (c) Electrochemical equivalent of ions  (d) Time for electrolysis		(a) 40.65 gm	(b) 4.065 <i>gm</i>			. ,			` '		
In an electroplating experiment $m$ $g$ of silver is deposited, when 4 amperes of current flows for 2 minutes. The amount (in $gms$ ) of silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991]  (a) $4m$ (b) $m/2$ (c) Electrochemical equivalent of ions (d) Time for electrolysis		(c) 0.4065 gm	(d) 65.04 <i>gm</i>		25		_	a fuead cale	` '		danosit on an
amperes of current flows for 2 minutes. The amount (in gms) of silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991]  (a) 4m  (b) m/2  (c) Electrochemical equivalent of ions  Time for electrolysis		In an electroplating ex	periment $m \ g$ of silver is	deposited, when 4	43.		,		uie	weight of the	•
silver deposited by 6 amperes of current flowing for 40 seconds will be [MNR 1991] (a) 4m (b) m/2 (c) Electrochemical equivalent of ions Time for electrolysis		amperes of current flo	ws for 2 minutes. The amo	ount (in gms) of				•			[=:
be [MNR 1991] (c) Electrochemical equivalent of ions (a) $4m$ (b) $m/2$ (d) Time for electrolysis		silver deposited by 6 a	mperes of current flowing f	or 40 seconds will			•				
(a) $4m$ (b) $m/2$ (d) Time for electrolysis		be [MNR 1991]						•	of ion	s	
		(a) 4 <i>m</i>	(b) $m/2$			( )		•	J. 1011	-	
(c) $m/4$ (d) $2m$ <b>26.</b> Faraday's laws of electrolysis will fail when [NCERT 1971]		(c) m/4	(d) 2 <i>m</i>		26	. ,		•	l fail v	vhen	[NCFRT 1071]

[MP PMT 1992]

27.

(a) Temperature is increased

(b) Inert electrodes are used

(d) In none of these cases

discharge at the electrode is

(a) W = ZQ

(c)  $W = \frac{Z}{F} It$ 

(c) A mixture of electrolytes is used

According to the first law of Faraday, the weight of a substance

(b) W = eF

(d) W = ZI

On passing 3 ampere of electricity for 50 minutes, 1.8 gram metal

The desired amount of charge for obtaining one mole of  $\ensuremath{\mathit{Al}}$  from

(b) 25.8

(d) 30.7

(b) 96500 C

deposits. The equivalent mass of metal is



28.		ectricity is passed through a solution of		(a) $9.65 \times 10^4$ Coulom	b (b) $6.28 \times 10^{18}$ Coulomb	<u></u>
		t of $Ca^{2+}$ metal deposited at the cathode		(c) $1.6 \times 10^{-19}$ Coulom		
	is [ <b>BHU 1996</b> ]					
	(a) 0.2 <i>gm</i>	(b) 0.4 <i>gm</i>	40.		passed through acidulated water 112	
	(c) 0.6 <i>gm</i>	(d) 0.8 gm		current passed, in amperes is	ect at the cathode in 965 <i>seconds</i> .	The
29.	•	for 5 hours through a molten metal salt		current passed, in amperes is	MNR 1991; UPSEAT	2001]
	metal in the metal salt is	At. wt. =177). The oxidation state of the [KCET 1996]		(a) 1.0	(b) 0.5	,
	(a) +1	(b) + 2		(c) 0.1	(d) 2.0	
	(c) + 3	(d) + 4	41.	How much chlorine will be	liberated on passing one ampere cu	rrent
30.	` '	ium will be deposited from a solution of		for 30 <i>minutes</i> through <i>NaC</i>	Cl solution [BVP 2003]	
		5 milliamperes flowing for 60 seconds		(a) 0.66 <i>mole</i> [ <b>BHU 1999</b> ]	(b) 0.33 <i>mole</i>	
				(c) 0.66 gm	(d) 0.33 gm	
	(a) $4.68 \times 10^{18}$	(b) $4.68 \times 10^{15}$	42.		volved in redox reactions when a Far	
	(c) $4.68 \times 10^{12}$	(d) $4.68 \times 10^9$			gh an electrolyte in solution is[NCEF	1 1982
31.	On passing 0.5 faraday of	electricity through $NaCl$ , the amount of		(a) $6 \times 10^{23}$	(b) $6 \times 10^{-23}$	
	Cl deposited on cathode	is		(c) 96500	(d) $8 \times 10^{19}$	
		[BHU 1997; RPET 1999]	43.	Coulomb is equal to	[Orissa JEE	2002]
	(a) 35.5 <i>gm</i>	(b) 17.75 <i>gm</i>		(a) ampere × second	(b) ampere × minute	
	(c) 71 <i>gm</i>	(d) 142 <i>gm</i>		(c) watt × second	(d) $volt \times second$	
32.	What is the amount of ch	lorine evolved when 2 amperes of current	4.4		· /	
	is passed for 30 minutes in	n an aqueous solution of NaCl [BHU 1998; A	411M <del>311</del> 999	9] The chergy required to relea	[Orissa JEE	20021
	(a) 66 <i>g</i> (c) 33 <i>g</i>	(b) 1.32 <i>g</i> (d) 99 <i>g</i>		(a) + 54.4 <i>eV</i>	(b) - 13.6 eV	
	.,	. , •		(c) $+ 27.2 \ eV$	(d) Cannot be predicted	
33.		ough $KCl$ solution, $19.5 g$ of potassium	45.	Faraday's laws of electrolysis	•	1983]
		quantity of electricity is passed through a		(a) Atomic number of cation	on	-
	is [EAMCET 1997]	oride, the amount of aluminium deposited		(b) Atomic number of anio	on	
	(a) 4.5 g	(b) 9.0 g		(c) Equivalent weight of th	e electrolyte	
	(c) 13.5 g	(d) 27 g		(d) Speed of the cation		
	(e) None is correct	( ) 5	46.		electrode decomposition of one	
34.	` '	raday's states that mass deposited on		equivalent of a substance is	[IIT 1984; KCET	1992]
	electrode is proportional t	•		(a) One ampere per second		
	(a) $m \propto I^2$	(b) $m \propto Q$		(b) 96500 coulombs per so		
	,	~ ~ ~		(c) One ampere for one ho		
	(c) $m \propto Q^2$	(d) None of these	457	(d) Charge on one mole of	electrons ssing per second through a cross-se	ation
35.		or two hour through a solution of an acid	47.	•		
		gen at NTP at anode. What will be the		of copper wire carrying 1	0 <sup>-6</sup> <i>amperes</i> of current per secon [EAMCET	
		ited at the cathode by the same current solution of copper sulphate for the same			•	1905]
	time [BVP 2003]	solution of copper surpliate for the sallie		(a) $1.6 \times 10^{-19}$	(b) $6 \times 10^{-35}$	
	(a) 16 g	(b) 63 g		(c) $6 \times 10^{-16}$	(d) $6 \times 10^{12}$	
	(c) 31.5 g	(d) 8 g	48.		ontaining acidified ferrous chloride	_
36.	· · · · · · · · · · · · · · · · · · ·	is 20% oxygen by weight. Its equivalent			ride are connected in series. The rate	_
	weight is	[Pb. PMT 2000]		through the cells will be	n the two cells when electricity is p	assea
	(a) 40	(b) 64		anough the cens will be	[CPMT	19891
	(c) 72	(d) 32		(a) 3:1	(b) 2:1	,
37.	On the basis of the i	nformation available from the reaction		(c) 1:1	(d) 3:2	
	$\frac{4}{2}Al + O_2 \rightarrow \frac{2}{2}Al_2O_2$	$\Delta G = -827 k J mol^{-1} \qquad \text{of} \qquad O_2 \qquad \text{the}$	49.	```	electricity is passed through a co	pper
	3				nt of copper deposited will be[MP PA	
	minimum <i>emf</i> required to	carry out an electrolysis of $Al_2O_3$ is $(F$		(a) 0.25 <i>mol</i>	(b) 0.50 <i>mol</i>	
	$= 96500 C \ mol^{-1})$	[CBSE PMT 2003]		(c) 1.00 <i>mol</i>	(d) 2.00 <i>mol</i>	
	(a) 8.56 V	(b) 2.14 V	50.	During electrolysis of fu	sed aluminium chloride 0.9 gm	of
	(c) 4.28 V	(d) 6.42 <i>V</i>			on the cathode. The volume of chl	
38.		of a solution of <i>AgNO</i> , 9650 <i>coulombs</i> of		liberated at the anode will b		
		e electroplating bath, the mass of silver		(a) 2.24 <i>litres</i>	(b) 11.2 <i>litres</i>	
	deposited in the cathode v			(c) 1.12 <i>litres</i>	(d) 5.6 <i>litres</i>	
	( )	[AIEEE 2003]	51.	Faraday has the dimensions	( ) -	1995]
	(a) 1.08 g	(b) 10.8 g	J	(a) Coulombs		[دور.
	(c) 21.6 g	(d) 108 g		(b) Coulomb equivalent		
39.	Total charge on 1 mole of a	monovalent metal ion is equal to		•	t	
<b>39</b> .	rotal charge on I mole of a	monovalent metal ion is equal to		(c) Coulomb per equivalen	t	

[DPMT 2001]

(d) Coulomb per degree Kelvin



52.		ivalent weight of silver deposite on				[AMl	J 1983; AFMC 19	89; MP PET 2001]
	cathode is	[Roorkee 1995]		(a)	$95500 \ C \ mol^{-1}$	(b)	96550 C n	$iol^{-1}$
	(a) $9.65 \times 10^7 C$	(b) $9.65 \times 10^4 C$		(c)	$96500 \ C \ mol^{-1}$	(d)	98500 C n	$iol^{-1}$
	(c) $9.65 \times 10^3 C$	(d) $9.65 \times 10^5 C$	63.	( )				
53.	96500 <i>coulombs</i> of electric curre	ent liberates from $CuSO_4$ solution	[MP PMT 199	95]	element is	aca to .		38; MP PMT 1997]
	(a) 63.5 <i>gm Cu</i>	(b) 31.75 <i>gm Cu</i>		(a)	48250 Faradays	(b)	48250 Coulo	mbs
	(c) 96500 gm Cu	(d) 100 gm Cu		. ,	193000 Faradays		193000 <i>Coul</i>	
54.		$np$ was passed through $CuSO_4$	64.		number of coulombs requer is	iired for	•	n of 107.870 <i>g</i> of <b>1P PET/PMT 1998</b> ]
	is	is. The amount of copper deposited		(a)	96,500	` ,	48,250	
	(Atomic weight of $Cu = 63.5$ )			( )	1,93,000	( )	10,000	
	(1 faraday = 96500 coulombs)		65.		en one of ampere current		or 1 sec throu	gh a conductor,
	( )	[EAMCET 1989; MP PET 1994]		this	quantity of electricity is k	nown as		[MD DMT recel
	(a) 0.3175 <i>g</i> (c) 0.635 <i>g</i>	(b) 3.175 <i>g</i> (d) 6.35 <i>g</i>		(2)	Faraday	(b)	Coulomb	[MP PMT 1993]
55.	.,	ty is passed through an aqueous	1	(c)	E.M.F.	(d)	Ohm	
00.	• •	c salt solution connected in series.		` '	mass deposited at an elect	` '		ional to
		s $1.08gm$ , the amount of copper			mass deposited at an elect		,	87; MP PET 2000]
	deposited is (atomic weight of C			(a)	At[ <b>EAMCET</b> ig <b>986</b> ]	-	Equivalent w	-
	(a) 0.6454 g	(b) 6.354 g		(c)	Molecular weight	(d)	Atomic numl	per
	(c) 0.3177 g	(d) 3.177 g	67.	Fro	m the solution of which of	the follo	owing one fara	day of electricity
56.	•	uired to deposit 1 <i>gm</i> atom of		will	liberate one gram atom of			
	be (where <i>N</i> is Avogadro's numb	solution of aluminium chloride will er)			•			2000; AFMC 2000]
	·	[AllMS 1992]		(a)	NaCl	(b)	$BaCl_2$	
	(a) 1 <i>N</i>	(b) 2 N		(c)	$CuSO_4$	(d)	$AlCl_3$	
57.	(c) 3 N  Three faradays of electricity are	(d) 4 ${\it N}$ e passed through molten $Al_2O_3$ ,	68.		electrolysis, 1 mole of a ten salt by	luminiur	n will be dep	oosited from its [MH CET 2000]
	aqueous solution of $\ensuremath{\textit{CuSO}_4}$ are	nd molten $NaCl$ taken in different		(a)	3 moles of electrons	(b)	4 moles of el	ectrons
	electrolytic cells. The amount of	Al, Cu and Na deposited at the	!	(c)	2 moles of electrons	(d)	1 mole of elec	trons
	cathodes will be in the ratio of	[BHU 1990]	69. 		atomic weight of $Fe$ is $Cl_3$ solution by passing 0.6			
	(a) 1 mole : 2 mole : 3 mole							[MH CET 2000]
	(b) 3 mole : 2 mole : 1mole (c) 1 mole : 1.5 mole : 3 mole			(a)	5.6 g	(b)	11.2 g	
	(d) 1.5 mole : 2 mole : 3 mole			(c)	22.4 g	(d)	33.6 g	
58.	An electrolytic cell contains a	solution of $Ag_2SO_4$ and have	70.	2.5	F of electricity are passe	ed throu	igh a <i>CuSO</i>	solution. The
	platinum electrodes. A current is	passed until 1.6 $\mathit{gm}$ of $O_2$ has been of silver deposited at cathode would		nun	nber of <i>gm</i> equivalent of <i>C</i>	•		is 82; MP PMT 2001]
	be	[CPMT 1971]		(a)	Zero	(b)	1.25	
	(a) 107.88 <i>gm</i>	(b) 1.6 <i>gm</i>			2.5	` '	5.0	
	(c) 0.8 gm	(d) 21.60 <i>gm</i>	71.		equivalent weight of a ce ght of its oxide is	rtain triv	alent element	
59.	The aqueous solution of which passing electric current	of the following decomposes on [EAMCET 1973]		'	152	(b)	56	[KCET 2003]
	(a) Canesugar	(b) Urea	l	. ,	168	(d)	68	
	(c) Methanol	(d) Potassium iodide	72.	Silv	er is removed electrically	from 2	00 <i>ml</i> of a 0	.1 N solution of
60.	The number of Faradays needed	d to reduce 4 gram equivalents of	f		$2NO_3$ by a current of 0			
	$Cu^{++}$ to $Cu$ metal will be	[BHU 1981]			nove half of the silver from			[AMU 1999]
	(a) 1	(b) 2		( )	16 <i>sec</i> 100 <i>sec</i>		96.5 <i>sec</i> 10 <i>sec</i>	
	(c) 1/2	(d) 4	. <b>73.</b>	. ,	order to separate oxygen f	` '		O the required
61.	,	d through the solution of			ntity of coulomb would be		T 1999]	o the required
		deposited. The number of Faraday	,		$1.93 \times 10^5$	=	$9.6 \times 10^4$	
	must be	ERT 1974; MP PET 1992; MP PMT 1994]		. ,	1.8		3.2	
	(a) 0.50	(b) 1.00	74.		current of $0.25A$ is passed	. ,		lution placed in
	(c) 1.50	(d) 2.00			ameter for 45 <i>minutes</i> . Th			
62.	The value of one Faraday is			is (	At weight of $Cu = 63.6$ )			<b>-</b>
								[BHU 2001]



[CPMT 1982;Pb.CET 2003]

	(c) 0.25 g	(d) 0.30 g		(a) Kohlraush	(b) Faraday
75.	Faraday constant	[KCET (Med.) 2001]		(c) Nernst	(d) Berthelot
	(a) Is a numerical constant		88.	` '	um will be deposited from a solution of
	(b) Depends on equivalent		00.		
	(c) Depends upon the current	nt passed		CuCi <sub>2</sub> by a current 0.23 7	nA following for 60 seconds
	(d) Depends on the number	of electrons		(a) $4.68 \times 10^{18}$	(b) $4.68 \times 10^{15}$
76.	If 0.5 amp current is passed t	hrough acidified silver nitrate solution		(c) $4.68 \times 10^{12}$	(d) $4.68 \times 10^9$
	for 10 minutes. The mass of s	ilver deposited on cathode, is (eq. wt.	_	` '	` '
	of silver nitrate = 108)	[AFMC 2001]	89.		required to reduce 12.3 g of nitrobenzene
	(a) 0.235 g	(b) 0.336 g		to aniline	[UPSEAT 2003]
	(c) 0.536 g	(d) 0.636 g		(a) 115800 <i>C</i>	(b) 5790 <i>C</i>
77.	The unit for the electric curren	nt is [KCET (Med.) 2001]		(c) 28950 C	(d) 57900 <i>C</i>
	(a) <i>Ohm</i>	(b) Volt	90.	During the process of ele-	ctrolytic refining of copper, some metals
	(c) Ampere	(d) Coulomb		present as impurity settle a	
78.	The quantity of electricity i	required to liberate $112 cm^3$ of			[AIEEE 2005]
<b>7</b> 0.	hydrogen at STP from acidified			(a) Sn and Ag	(b) $Pb$ and $Zn$
	nydrogen at 317 from acidillet	[KCET (Med.) 2001]		(c) Ag and Au	(d) Fe and Ni
	(a) 0.1 <i>Faraday</i>	(b) 1 Faraday	01	( ) 2	· /
	(c) 965 <i>Coulomb</i>	(d) 96500 <i>Coulomb</i>	91.		n a zinc bar weighing 50 <i>g</i> and 1.0 <i>litre</i> , 1.0 <i>M</i> ,
79.	( )	hest resistance during the passage of			g would the cell run, assuming it delivers a
19.	current	[BHU 2001]		steady current of 1.0 ampere	
		(b) 2 N NaCl			[Roorkee 2000]
	(a) 0.05 <i>N NaCl</i>			(a) 48 <i>hrs</i>	(b) 41 <i>hrs</i>
_	(c) 0.1 <i>N NaCl</i>	(d) 1 N NaCl		(c) 21 <i>hrs</i>	(d) 1 <i>hr</i>
80.		on concentrated nitric acid. The copper	92.	On passing electric curren	t through molten aluminium chloride, 11.2
	equivalent weight of copper is	neating gave 5 $g$ of its oxide. The [KCET 2004]	_	· -	d at NTP at anode. The quantity of
				aluminium deposited at cat	
	(a) 23	(b) 32		(a) 9 g	(b) 18 g
٥.	(c) 12	(d) 20			
81.		sited by passing 241.25 coulomb of		(c) 27 g	(d) 36 g
	current through silver nitrate s		93.		ed through silver voltameter connected to
	(a) 2.7 g	(b) 2.7 mg			eathode of the silver voltameter weighed of the electrolysis. The volume of oxygen
_	(c) 0.27 g	(d) 0.54 g		evolved at STP is	of the electrolysis. The volume of oxygen
82.		passed through acidulated water,		everved de 511 ib	[Kerala (Med.) 2003]
	$O_2$ evolved is	[MHCET 2004]			
	(a) $11.2 dm^3$	(b) $5.6 dm^3$		(a) 56 <i>cm</i> <sup>3</sup>	(b) 550 $cm^3$
	(a) 11.2 um			(c) 5.6 cm <sup>3</sup>	(d) 11.2 $cm^3$
	(c) $22.4 \ dm^3$	(d) $1.0 \ dm^3$			(a) <u>.</u>
83.	Charge required to liberate 11.5	a codium ic		(e) 22.4 <i>cm</i> <sup>3</sup>	
03.	Charge required to inserate in.	_	94.	During electrolysis of aque	ous $NaOH, 4g$ of $O_2$ gas is liberated
	() 27 5	[AllMS 1992; DCE 2002]			
	(a) 0.5 F	(b) 0.1 <i>F</i>		at $NTP$ at anode, $H_2$ ga	
	(c) 1.5 F	(d) 96500 <i>coulombs</i>			[CBSE PMT 1998]
84.		ne Faraday of electrical energy would		(a) 2.8 <i>litres</i>	(b) 5.6 <i>litres</i>
	evolve	[DCE 2004]		(c) 11.2 <i>litres</i>	(d) 22.4 <i>litres</i>
	(a) One mole of oxygen	(b) One $g$ atom of oxygen			
	(c) $8 g$ of oxygen	(d) 22.4 <i>litres</i> of oxygen		Conductor a	and conductance
85.	In a galvanic cell, the electrons	flow from [KCET 2004]			
	(a) Anode to cathode through	h the solution	1.	Which one of the following	statements is correct
	(b) Cathode to anode through	h the solution	••	winer one of the following	[MP PET 1997]
	(c) Anode to cathode through			() m :1.: 1	
	(d) Cathode to anode through			(a) The oxidation number	of oxygen in $KO_2$ is zero
86.	. ,			(b) The specific conducts	ance of an electrolyte solution decreases
<b>60.</b>		through silver nitrate solution using ver was found to be deposited on the		with increase in diluti	on
		of electricity is passed through copper		(c) $Sn^{2+}$ oxidises $Fe^{3+}$	
		per electrodes, the weight of copper		( )	1 . 1
	deposited on the cathode is	5 Tr		(d) $Zn/ZnSO_4$ is a ref	erence electrode
		[Kerala PMT 2004]	2.	In infinite dilutions, the ear	uivalent conductances of $Ba^{2+}$ and $Cl^{-}$
	(a) 6.4 g	(b) 2.3 g		•	
	(c) 12.8 g	(d) 1.6 g			$eqvt^{-1}$ . The equivalent conductivity
		(4) 1.0 5		of $BaCl_2$ at indefinite dilu	tion is [CBSE 2000]
0-	(e) 3.2 g			(a) 101.5	(b) 139.5
87.	The law of electrolysis were pr	oposea by			

(a) 0.20 g

(b) 0.22 g

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	(c) 203.5 (d) 279.5		(c) ohm cm² (gm equivalent)
3.	The factor which is not affecting the conductivity of any solution is		
	(a) Dilution (b) Nature of electrolyte		(d) $S cm^{-2}$
	(c) Temperature (d) None of these	14.	It has been observed that gaseous hydrogen chloride is a very poor
4.	Specific conductance of 0.1 <i>m</i> nitric acid is		conductor of electricity but a solution of hydrogen chloride gas in water is a good conductor of electricity. This is due to the fact that[N
	$6.3 \times 10^{-2} \ ohm^{-1} cm^{-1}$ . The molar conductance of solution is		(a) Water is good conductor of electricity
	[Kerala (Med.) 2003]		(b) Hydrogen chloride gas in water solution ionizes
			(c) A gas is non-conductor but a liquid conducts electricity
			(d) Gas does not obey Ohm's law whereas solution does
	(c) $100 \ ohm^{-1}cm^2mole^{-1}$ (d) $6300 \ ohm^{-1}cm^2mole^{-1}$	15.	Electrolytic conduction differs from metallic conduction in that in
	(e) $63.0 \ ohm^{-1}cm^2mole^{-1}$		the case of electrolytic conduction
5.	The conductivity of strong electrolyte is [CPMT 2003]		[KCET 1987; Bihar CEE 1992]
	(a) Increase on dilution slightly		(a) The resistance increases with increasing temperature
	(b) Decrease on dilution		(b) The resistance decreases with increasing temperature
	(c) Does not change with dilution		(c) The flow of current does not generate heat
	(d) Depend upon density of electrolytes itself		(d) The resistance is independent of the length of the conductor
6.	If V is the specific resistance of the solution and M is the molarity of	16.	The electrolytic conductance is a direct measure of
	the solution, the molar conductivity of the solution is given by [Kurukshe	etra CEE 2	[KCET 1990; CPMT 2003] (a) Resistance (b) Potential
	1000X $1000$		(d) Resistance (b) Potential (c) Concentration (d) Dissociation
	(a) $\frac{1}{M}$ (b) $\frac{1}{MX}$	17.	Conductivity of a strong electrolyte [KCET 1993]
	400014	.,.	(a) Increases on dilution
	(c) $\frac{1000M}{X}$ (d) $\frac{MX}{1000}$		(b) Does not change considerably on dilution
_	X 1000		(c) Decreases on dilution
7.	Conductivity (unit Siemen's) is directly proportional to area of the vessel and the concentration of the solution in it and is inversely		(d) Depends on density
	proportional to the length of the vessel then the unit of the constant	18.	Which of the following statements is not applicable to electrolytic
	of proportionality is [AIEEE 2002]		conductors [AIIMS 1991]
	(a) $Sm \ mol^{-1}$ (b) $Sm^2 \ mol^{-1}$		(a) New products show up at the electrodes
	· · ·		(b) lons are responsible for carrying the current
	(c) $S^{-2}m^2 \ mol$ (d) $S^2m^2 \ mol^{-2}$		(c) Show a positive temperature coefficient for conductance
8.	If one end of a piece of a metal is heated, the other end becomes hot		(d) A single stream of electrons flows from cathode to anode
	after some time. This is due to	19.	Which one is not a conductor of electricity [RPET 1999]
	[CBSE PMT 1995]		(a) NaCl (aqueous) (b) NaCl (solid)
	(a) Energised electrons moving to the other part of the metal		(c) NaCl (molten) (d) Ag metal
	(b) Resistance of the metal	20.	Solid sodium chloride is bad conductor of electricity because
	(c) Mobility of atoms in the metal		(a) It contains only molecules
	(d) Minor perturbation in the energy of atoms		(b) It does not possess ions
9.	Conductivity of a solution is directly proportional to		(c) The ions present in it are not free to move
	[KCET 1984]		(d) It does not contain free molecules
	(a) Dilution (b) Number of ions	21.	Which of the following is a poor conductor of electricity
10	(c) Current density (d) Volume of the solution		[EAMCET 1992]
10.	The increase in equivalent conductance of an electrolyte solution with dilution is due to the increase in		(a) $CH_3COONa$ (b) $C_2H_5OH$
	[MP PMT 1996]		(c) NaCl (d) KOH
	(a) lonic attraction	22.	The molar conductivity is maximum for the solution of
	(b) Molecular attraction		concentration [DCE 2002]
	(c) Degree of association of the electrolyte		(a) 0.001 M (b) 0.005 M
	(d) Degree of ionisation of the electrolyte	22	(c) 0.002 M (d) 0.004 M  The unit of molar conductivity is
11.	Which of the following conducts electricity [AFMC 1995]	23.	The unit of molar conductivity is [DCE 2002]
	(a) Fused $NaCl$ (b) $CO_2$		(a) $\Omega^{-1}cm^{-2}mol^{-1}$ (b) $\Omega \ cm^{-2}mol^{-1}$
	( ) 2		(c) $\Omega^{-1}cm^2mol^{-1}$ (d) $\Omega cm^2mol$
	( )	24.	The highest electrical conductivity of the following aqueous solutions
12.	Which of the following shows electrical conduction		is of [AIEEE 2005]

[CBSE PMT 1999; AlIMS 1999]

[CPMT 1999; BCECE 2005]

(b) Graphite

(d) Sodium

(a) Potassium

(c) Diamond

(a) ohm cm

The unit of equivalent conductivity is

(b)  $ohm^{-1}cm^{2}(gm \ equivalent)^{-1}$ 

(a) 0.1 M acetic acid

(c) 0.1 M fluoroacetic acid

(a)  $10 ohm^{-1} cm^2 gm eq^{-1}$ 

(b) 0.1 M chloroacetic acid

Given  $l/a = 0.5 cm^{-1}$ , R = 50 ohm, N = 1.0. The equivalent

(c)  $300 \, ohm^{-1} cm^2 \, gmeq^{-1}$  (d)  $100 ohm^{-1} cm^2 \, gmeq^{-1}$ 

conductance of the electrolytic cell is [Orissa JEE 2005]

(d) 0.1 M difluoroacetic acid

(b)  $20 ohm^{-1} cm^2 gm eq^{-1}$ 



- If equivalent conductance of 1M benzoic acid is  $12.8 ohm^{-1} cm^2$ 26. and if the conductance of benzoate ion and  $H^{\scriptscriptstyle +}$  ion are 42 and 288.42 ohm<sup>-1</sup> cm<sup>2</sup> respectively. its degree of dissociation is
  - (a) 39%
- (b) 3.9%
- (c) 0.35%
- (d) 0.039%
- The unit  $ohm^{-1}$  is used for

[] & K 2005]

- (a) Molar conductivity
- (b) Equivalent conductivity
- (c) Specific conductivity
- (d) Conductivity

#### Cell constant and Electrochemical Cells

- When electric current is passed through a cell having an electrolyte, the positive ions move towards the cathode and the negative ions towards the anode. If the cathode is pulled out of the solution [AIIMS 1980]10.
  - (a) The positive and negative ions will move towards the anode
  - (b) The positive ions will start moving towards the anode, the negative ions will stop moving
  - The negative ions will continue to move towards the anode and the positive ions will stop moving
  - (d) The positive and negative ions will start moving randomly
- If the half cell reaction  $A + e^- \rightarrow A^-$  has a large negative 2. reduction potential, it follows that

#### [MNR 1992; UPSEAT 2000, 02]

- (a) A is readily reduced
- (b) A is readily oxidised
- (c)  $A^-$  is readily reduced
- (d)  $A^-$  is readily oxidised
- Mark the false statement 3.

- (a) A salt bridge is used to eliminate liquid junction potential
- (b) The Gibbs free energy change,  $\Delta G$  is related with electromotive force (E), as  $\Delta G = -nFE$
- (c) Nernst equation for single electrode  $E = E^o - \frac{RT}{nF} \ln a_{M^{n+}}$
- (d) The efficiency of a hydrogen oxygen fuel cell is 23%
- The specific conductance of a 0.1 N KCl solution at  $23^{\circ}C$  is  $0.012\,ohm^{-1}\,cm^{-1}$  . The resistance of cell containing the solution at the same temperature was found to be 55 ohm. The cell constant will be

#### [CBSE PMT 1999, 2000; KCET 2001]

- (a)  $0.142 \, cm^{-1}$
- (b)  $0.66 \, cm^{-1}$
- (c)  $0.918 \, cm^{-1}$
- (d)  $1.12 \, cm^{-1}$
- Which of the following reactions occurs at the cathode of a common dry cell [NCERT 1978]
  - (a)  $Mn \to Mn^{2+} + 2e^{-}$
  - (b)  $2MnO_2 + Zn^{2+} + 2e^- \rightarrow ZnMn_2O_4$
  - (c)  $2ZnO_2 + Mn^{2+} + 2e^- \rightarrow MnZn_2O_A$
  - (d)  $Zn \rightarrow Zn^{2+} + 2e^{-}$
- In Cu Zn cell 6.

[BHU 1981]

- (a) Reduction occurs at the copper cathode
- (b) Oxidation occurs at the copper cathode
- (c) Reduction occurs at the anode
- (d) Chemical energy is converted to light energy
- Which of the following reaction is used to make a fuel cell 7.

[AIIMS 2003]

- (a)  $Cd(s) + 2Ni(OH)_3(s) \rightarrow CdO(s) + 2Ni(OH) + H_2O(l)$
- $Pb(s) + PbO_2(s) + 2H_2SO_4(aq) \rightarrow 2PbSO_4(s) + 2H_2O(l)$
- $2H_2(g) + O_2(g) \rightarrow 2H_2O(l)$

- $2Fe(s) + O_2(g) + 4H^+(aq) \rightarrow 2Fe^{2+}(aq) + 2H_2O(l)$
- 8. When lead storage battery is charged

[MP PET 2003]

- P[DPMTs 2005] lved
- (b)  $H_2SO_4$  is regenerated
- (c)  $PbSO_4$  is deposited on lead electrode
- (d) Lead is deposited on lead electrode
- 9. When lead storage battery is charged

[MP PET 1993; MP PMT 2000]

- (a) Lead dioxide dissolves
- (b) Sulphuric acid is regenerated
- The lead electrode becomes coated with lead sulphate
- (d) The amount of sulphuric acid decreases

The electrolytic decomposition of dilute sulphonic acid with platinum electrode in cathodic reaction is

[MNR 1988; UPSEAT 1999, 2002]

- (a) Oxidation
- (b) Reduction
- (c) Oxidation and reduction both
- (d) Neutralisation
- Which colourless gas evolves, when  $NH_{4}Cl$  reacts with zinc in a 11. dry cell battery [Orissa JEE 2003]
  - (a)  $NH_4$
- (b)  $N_2$
- (c)  $H_2$
- (d)  $Cl_2$
- Which of the substances Na, Hg, S, Pt and graphite can be used as electrodes in electrolytic cells having aqueous solutions
  - (a) Na, Pt and graphite
- (b) Na and Hg
- (c) Pt and graphite only
- (d) Na and S only
- In electrolysis of dilute  $H_2SO_4$  using platinum electrodes 13.

#### [DPMT 1983; IIT 1983; Kurukshetra CET 2002; AFMC 2005]

- (a)  $H_2$  is evolved at cathode
- (b)  $NH_3$  is produced at anode
- (c)  $Cl_2$  is obtained at cathode
- (d)  $O_2$  is produced
- For cell reaction,  $Zn + Cu^{2+} \rightarrow Zn^{2+} + Cu$ , cell representation is 14.
  - (a)  $Zn \mid Zn \mid Cu \mid Cu$
- (b)  $Cu \mid Cu \mid |Zn| \mid Zn$
- (c) Cu | Zn | Zn | Cu
- (d)  $Cu \mid Zn \mid Zn \mid Cu$
- Which one is not called a anode reaction from the following
  - (a)  $Cl^{-} \rightarrow \frac{1}{2} Cl_{2} + e^{-}$  (b)  $Cu \rightarrow Cu^{++} + 2e^{-}$
  - (c)  $Hg^+ \to Hg^{++} + e^-$
- (d)  $Zn^{2+} + 2e^- \rightarrow Zn$
- A cell from the following which converts electrical energy into 16. chemical energy
  - (a) Dry cell
- (b) Electrochemical cell
- (c) Electrolytic cell
- (d) None of these
- In the cell  $Zn \mid Zn^{2+} \mid Cu^{2+} \mid Cu$ , the negative electrode is 17.

[MP PMT 1995]

- (a) Cu
- (b)  $Cu^{2+}$
- (c) Zn
- (d)  $Zn^{2+}$
- Which of the following statements is correct? Galvanic cell converts[KCET 1991 18.
  - (a) Chemical energy into electrical energy
  - (b) Electrical energy into chemical energy
  - Metal from its elemental state to the combined state



- (d) Electrolyte into individual ions
- Hydrogen-oxygen fuel cells are used in space-craft to supply [MP PMT 1993; MP PE901999] 19.
  - (a) Power for heat and light
  - (b) Power for pressure
  - Oxygen
  - (d) Water
- The standard cell potential of  $Zn|Zn^{2+}_{(aq)}||Cu^{2+}_{(aq)}|Cu$  cell is 20. 1.10 V. The maximum work obtained by this cell will be
  - (a) 106.15 kl
- (b) -212.30 kI
- (c) -318.45 k/
- (d) 424.60 kJ
- The relationship between standard reduction potential of cell and 21. equilibrium constant is shown by
  - (a)  $E_{cell}^{0} = \frac{n}{0.059} \log K_c$  (b)  $E_{cell}^{0} = \frac{0.059}{n} \log K_c$
  - (c)  $E_{cell}^0 = 0.059 n \log K_c$  (d)  $E_{cell}^0 = \frac{\log K_c}{n}$
- Consider the Galvanic cell  $Zn^{\Theta} | ZnSO_4 | CuSO_4 | Cu^{\oplus}$  the 22. reaction at cathode is [AMU 2000]
  - (a)  $Zn^{2^+} + 2e^- \rightarrow Zn$
  - (b)  $Cu^{2+} + 2e^{-} \rightarrow Cu$
  - (c)  $Cu^{2+} + Zn \rightarrow Cu + Zn^{2+}$
  - (d)  $Zn^{2+} + Cu \rightarrow Zn + Cu^{2+}$
- The cell reaction  $Cu + 2Ag^+ \rightarrow Cu^{+2} + Ag$  is best represented 23.
  - (a)  $Cu_{(s)} | Cu^{+2}_{(aq)} | | Ag^{+}_{(aq)} | Ag_{(s)}$
  - (b)  $Pt | Cu^{+2} | | Ag^{+}_{(aq)} | Ag_{(s)}$
  - (c)  $Cu^{+2} | Cu | | Pt | Ag$
  - (d) None of the above representations
- $Zn_{(s)} | Zn^{2+}_{(aq)} | | Cu^{2+}_{(aq)} | Cu_{(s)}$  is [Kerala (Engg.) 2002]
  - (a) Weston cell
- (b) Daniel cell
- (c) Calomel cell
- (d) Faraday cell
- (e) Standard cell
- The specific conductance of a solution is  $0.2 ext{ } ohm^{-1}cm^{-1}$  and 25. conductivity is 0.04  $ohm^{-1}$  . The cell constant would be
  - (a) 1  $cm^{-1}$
- (c) 5  $cm^{-1}$
- (d)  $0.2 \ cm^{-1}$
- If the conductance and specific conductance of a solution is one 26. then its cell constant would be [RPET 1999]

(b) Zero

(c) 0.5

- (d) 4
- Saturated solution of KNO3 is used to make 'salt-bridge' because 27.
  - (a) Velocity of  $K^+$  is greater than that of  $NO_3^-$
  - (b) Velocity of  $NO_3^-$  is greater than that of  $K^+$
  - (c) Velocities of both  $K^+$  and  $NO_3^-$  are nearly the same
  - (d) KNO3 is highly soluble in water
- In balancing the half reaction  $S_2O_3^{2-} \rightarrow S_{(s)}$  the number of 28. electrons that must be added is [DPMT 2000]
  - (a) 4 on the left
- (b) 3 on the right
- (c) 2 on the left
- (d) 2 on the right

- Which one of the following statement is true for a electrochemical [Pb. PMT 1999; KCET 1999]
  - (a)  $\dot{H}_2$  is cathode and Cu is anode
  - (b)  $H_2$  is anode and Cu is cathode
  - (c) Reduction occurs at  $H_2$  electrode
  - (d) Oxidation occurs at Cu electrode
- 30. In the reaction

[MP PET 2002] 
$$Ag^{+}(aq) \rightarrow Cu^{2+}(aq) + 2Ag(s)$$

The reduction half-cell reaction is [AIIMS 1997]

- (a)  $Cu + 2e^- \rightarrow Cu^{2-}$
- (b)  $Cu 2e^- \rightarrow Cu^{2+}$
- (c)  $Ag^+ + e^- \rightarrow Ag$
- (d)  $Ag e^- \rightarrow Ag^+$
- Which of the following statements about galvanic cell is incorrect[JIPMER 1997] 31.
  - (a) Anode is positive
  - (b) Oxidation occurs at the electrode with lower reduction potential
  - Cathode is positive
  - (d) Reduction occurs at cathode
- The molar conductances of NaCl, HCl and CH3COONa at 32. infinite dilution are 126.45, 426.16 and  $91 \, ohm^{-1} \, cm^2 \, mol^{-1}$ respectively. The molar conductance of CH3COOH at infinite [CBSE PMT 1997]
  - (a)  $201.28 \, ohm^{-1} \, cm^2 \, mol^{-1}$
  - (b)  $390.71 ohm^{-1} cm^2 mol^{-1}$
  - (c)  $698.28 \, ohm^{-1} \, cm^2 \, mol^{-1}$
  - (d)  $540.48 \, ohm^{-1} cm^2 \, mol^{-1}$
- The electrodes of a conductivity cell are 3 cm apart and have a 33. cross-sectional area of  $4 cm^2$ . The cell constant of the cell (in  $cm^{-1}$ ) is
  - (a)  $4 \times 3$
- (b) 4/3
- (d) 9/4
- The anode half-reaction occurring during the discharge of a lead storage battery is
  - (a)  $Pb(s) + SO_2 + O_2 \rightarrow PbSO_4(s)$
  - (b)  $Pb(s) + SO_4^{2-}(aq) = PbSO_4(s) + 2e^{-1}$
  - (c)  $PbO_2(s) + 4H^+(aq) + 2e^- + SO_4^{2-}(aq) \Rightarrow$

[RPET 1999]

 $PbSO_{4}(s) + 2H_{2}O$ 

- (d)  $Pb^{2+}(aq) + SO_4^{2-}(aq) \rightarrow PbSO_4(s)$
- The unit of cell constant is 35.

[MP PET 1996]

- (a)  $ohm^{-1}cm^{-1}$
- (b) ohm cm
- (d)  $cm^{-1}$
- 36. In dry coll the reaction which takes place at the zinc anode is

[MP PET 1996]

- (a)  $Zn^{2+} + 2e^- \rightarrow Zn(s)$
- (b)  $Zn(s) \rightarrow Zn^{2+} + 2e^{-}$
- (c)  $Mn^{2+} + 2e^- \rightarrow Mn(s)$
- (d)  $Mn(s) \to Mn^+ + e^- + 1.5V$
- The chemical reaction taking place at the anode of a cell is 37.

[MP PET 1996]

- (a) lonisation
- (b) Reduction



- Oxidation (d) Hydrolysis Create potential difference between the two electrodes Generate heat 38. Which of the following reactions occurs at the cathode during the (d) Remove adsorbed oxygen from electrode surfaces charging of a lead storage battery [Manipal MEE 1995; MP PET 2002]  $\lambda_{CICH_2COONa} = 224 \, ohm^{-1} cm^2 gmeg^{-1}$ ,  $Pb^{2+} + 2e^{-} \rightarrow Pb$  $\lambda_{NaCl} = 38.2 \, ohm^{-1} cm^2 gmeg^{-1}$ ,  $Pb^{2+} + SO_4^{2-} \rightarrow PbSO_4$  $\lambda_{HCI} = 203 \, ohm^{-1} cm^2 gmeq^{-1} \,,$ (c)  $Pb \rightarrow Pb^{2+} + 2e^{-}$ What is the value of  $\lambda_{CICH_2COOH}$ [JEE Orissa 2004] (d)  $PbSO_4 + 2H_2O \rightarrow 2PbO_2 + 4SO_4^{2-} + 2e^{-}$ (a)  $288.5 \text{ ohm}^{-1} \text{cm}^2 \text{gmeg}^{-1}$ A depolarizer used in dry cell batteries is 39. [NCERT 1981]  $289.5 \ ohm^{-1}cm^{2}gmeg^{-1}$ (a) Ammonium chloride (b) Manganese dioxide  $388.5 \ ohm^{-1}cm^{2}gmeg^{-1}$ (d) Sodium phosphate (c) Potassium hydroxide When a lead storage battery is discharged 40. (d)  $59.5 \ ohm^{-1}cm^2gmeg^{-1}$ [IIT 1987; MP PMT 2004] Which of the following statement is true for the electrochemical 50. (a)  $SO_2$  is evolved Daniel cell Electrons flow from copper electrode to zinc electrode (b) Lead sulphate is consumed Current flows from zinc electrode to copper electrode (c) Lead is formed (c) Cations move toward copper electrode which is cathode (d) Sulphuric acid is consumed Cations move toward zinc electrode In electroplating, the article to be electroplated serves as 41. Which of the following statement is true for an electrochemical cell 51. [AMU 1982, 83]  $H_2$  is cathode and Cu is anode (a) Cathode (b) Electrolyte  $H_2$  is anode and Cu is cathode (c) Anode (d) Conductor 42. The position of some metals in the electrochemical series in (c) Reduction occurs at  $H_2$  electrode electropositive character is (d) Oxidation occurs at Cu electrode Mg > Al > Zn > Cu > Ag. What will happen, if a copper spoon Which of the following statements are true for an fuel cells is used to stir a solution of aluminium nitrate [DPMT 2004] [NCERT 1977] They run till the reactants are active (a) The spoon will get coated with AlThey are free from pollution (b) An alloy of Cu and Al is formed They are more efficient The solution becomes blue All of the above There is no reaction For gold plating, the electrolyte used is [Pb.CET 2004] 53. 43. In a electrochemical cell [AFMC 1989]  $AuCl_3$ (b) HAuCl<sub>4</sub> (a) Potential energy changes into kinetic energy (c)  $k[Au(CN)_2]$ (d) None of these Kinetic energy changes into potential energy (c) Chemical energy changes into electrical energy The acid used in lead storage battery is [Pb.CET 2003] 54. Electrical energy changes into chemical energy (a)  $H_2SO_4$ (b)  $H_3PO_4$ In galvanic cell, the salt bridge is used to [MP PMT 2002] Complete the circuit (c) HCl (d)  $HNO_3$ (a) Reduce the electric resistance in the cell (b) At 25°C specific conductivity of a normal solution of KCl is 55. Separate cathode from anode (c) Carry salts for the chemical reaction [Pb.PMT 2004] If a strip of Cu metal is placed in a solution of ferrous sulphate [NCERT 1974; CPMT] 1977; MP PET 2000] 45 (b) 1.016 (a) Copper will precipitate out (c) 1.106 (d) 2.016 (b) Iron will precipitate out Which of the following is used widely in the manufacture of lead (c) 56. Copper will dissolve storage battery No reaction will take place Which of the following is not used to construct salt bridge (b) Lithium (a) Arsenic [MP PET 2003] Bismuth (d) Antimony (c) (a) CH3COOK (b) KCl

(d)  $KNO_3$ 

- The reference electrode is made by using [MP PMT 2002] 47.
- - (a)  $ZnCl_2$ (b)  $CuSO_4$
  - (c)  $HgCl_2$ (d)  $Hg_2Cl_2$
- In a hydrogen oxygen fuel cell, combustion of hydrogen occurs to 48.
  - (a) Produce high purity water

(c)  $NH_4NO_3$ 

- 0.002765 mho. The resistance of cell is 400 ohms. The cell constant
- [BHU 2004]
- The  $2AgCl(s) + H_2(g) \rightarrow$ 57. chemical reaction, 2HCl(aq) + 2Ag(s) taking place in a galvanic cell is represented [AIIMS 2005] by the notation
  - $Pt|H_2(g)$ , 1bar | 1M KCl(aq)/AgCl(s)|Ag(s)
  - Pranel Local, 1bar |  $1M \ HCl(aq)$  |  $1M \ Ag^+(aq)$  | Ag(s)

- $Pt(s)|H_2(g)$ , 1bar | 1M HCl(aq)|AgCl(s)|Ag(s)
- (d)  $Pt(s) \mid H_2(g)$ , 1bar | 1M  $HCl(aq) \mid Ag(s) \mid AgCl(s)$
- If the  $Zn^{2+}/Zn$  electrode is diluted to 100 times then the change 58.
  - (a) Increase of  $59 \, mV$
  - Decrease of 59 mV
  - Increase of  $29.5 \, mV$
  - (d) Decrease of  $29.5 \, mV$
- 59. If hydrogen electrode dipped in 2 solution of pH = 3 and pH = 6 and salt bridge is connected the e.m.f. of resulting cell is [DPMT 2005]
  - (a) 0.177 V
- (b) 0.3 V
- (c) 0.052 V
- 60. The tendency of an electrode to lose electrons is known as

[] & K 2005]

- (a) Electrode potential
- (b) Reduction potential
- (c) Oxidation potential
- (d) e.m.f.
- When electric current is supplied through an ionic hydride of fused 61. state, then [Kerala CET 2005]
  - (a) Hydrogen is obtained at anode
  - (b) Hydrogen is obtained at cathode
  - (c) No change
  - (d) Hydride ion moves towards cathode
  - (e) hydride ion present in solution

## Electrode potential, Ecel, Nernt equation and ECS

- The hydrogen electrode is dipped in a solution of pH = 3 at 1.  $25^{\circ}C$ . The potential of the cell would be (the value of 2.303RT/Fis 0.059 V [KCET 1993,2005]
  - (a) 0.177 V
- (b) -0.177 V
- (c) 0.087 V
- (d) 0.059 V
- The standard electrode potentials of  $Zn^{2+}/Zn$  and  $Ag^+/Ag$ 2. are -0.763 V and +0.799 V respectively. The standard potential of the cell is [KCET 1993]
  - (a) 1.56 V
- (b) 0.036 V
- (c) -1.562 V
- (d) 0.799 V

-0.762

-0.740

The standard reduction potentials at 298 K for the following half 3. reactions are given against each

$$Zn^{2+}(aq.) + 2e \Rightarrow Zn(s);$$

$$Cr^{3+}(aq) + 3e \Rightarrow Cr(s)$$
;

$$2H^{+}(aq) + 2e \Rightarrow H_{2}(g);$$
 0.00

$$Fe^{3+}(aq) + e \Rightarrow Fe^{2+}(aq);$$
 0.770

Which is the strongest reducing agent

[11T 1981; MP PET/PMT 1988; MP PMT 1989; MH CET 2001

- (a) Zn(s)
- (b) Cr(s)
- (c)  $H_2(g)$
- (d)  $Fe^{2+}(aq)$
- When Zn piece is kept in  $CuSO_4$  solution, the copper get precipitated due to standard potential of zinc is

[CPMT 1999]

- (a) > copper
- (b) < copper
- (c) > sulphate
- (d) < sulphate
- Which of the following metal does not react with the solution of copper sulphate [CPMT 1999]
  - (a) Mg
- (b) *Fe*
- Zn(c)

6.

7.

- (d) Ag
- solution containing one mole per litre of  $Cu(NO_3)_2$ ,  $AgNO_3$ ,  $Hg_2(NO_3)_2$  and  $Mg(NO_3)_2$ , is being electrolysed by using inert electrodes. The values of standard electrode potentials in volts (reduction potentials)  $Ag/Ag^{+} = +0.80, 2Hg/Hg_{2}^{2+} = +0.79, Cu/Cu^{2+} = +0.34,$

 $Mg/Mg^{2+} = -2.37$  with increasing voltage, the sequence of deposition of metals on the cathode will be

[IIT 1984; AMU 1999; Kerala PMT 2004]

- (a) Ag, Hg, Cu, Mg
- (b) Mg, Cu, Hg, Ag
- (c) Ag, Hg, Cu
- (d) Cu, Hg, Ag
- The standard reduction electrode potentials of four elements are

$$A = -0.250 V$$
  
 $C = -0.126 V$ 

$$B = -0.136 V$$

$$C = -0.126 V$$

$$D = -0.402 \ V$$

The element that displaces A from its compounds is

(a) B

- (b) C
- (c) D
- (d) None of these
- 8. The standard oxidation potential of zinc and silver in water at

$$Zn(s) \rightarrow Zn^{2+} + 2e^{-}; E = 0.76 V$$

$$Ag(s) \rightarrow Ag^{2+} + 2e^{-}; E = -0.80 V$$

Which of the following reactions actually take place

[NCERT 1983, 84; KCET 2003]

- (a)  $Zn(s) + 2Ag^+(aq) \rightarrow Zn^{++}(aq) + 2Ag(s)$
- (b)  $Zn^{++}(aq) + 2Ag(s) \rightarrow 2Ag^{+}(aq) + Zn(s)$
- (c)  $Zn(s) + Ag(s) \rightarrow Zn^{++}(aq) + Ag^{+}(aq)$
- (d)  $Zn^{++}(aq) + Ag^{+}(aq) \rightarrow Zn(s) + Ag(s)$
- Beryllium is placed above magnesium in the second group. Beryllium dust, therefore when added to MgCl2 solution will

[CPMT 1977]

- (a) Have no effect
- (b) Precipitate Mg metal
- (c) Precipitate MgO
- (d) Lead to dissolution of Be metal
- The name of equation showing relation between electrode potential (E) standard electrode potential  $(E^{o})$  and concentration of ions in solution is
  - (a) Kohlrausch's equation
- (b) Nernst's equation
- (c) Ohm's equation
- (d) Faraday's equation
- The correct representation of Nernst's equation is

(a) 
$$E_{M^{n+}/M} = E^{o}_{M^{n+}/M} + \frac{0.0591}{n} \log(M^{n+})$$

(b) 
$$E_{M^{n+}/M} = E_{M^{n+}/M}^{o} - \frac{0.0591}{n} \log(M^{n+})$$



- (c)  $E_{M^{n+}/M} = E^{o}_{M^{n+}/M} + \frac{n}{0.0591} \log(M^{n+})$
- (d) None of the above
- Standard electrode potential of NHE at 298 K is 12.
  - (a) 0.05 V
- (b) 0.1 V
- (c) 0.00 V
- (d) 0.11 V
- When a copper wire is placed in a solution of  $AgNO_3$ , the solution 13. acquires blue colour. This is due to the formation of

[Roorkee 1989]

- (a)  $Cu^{2+}$  ions
- (b)  $Cu^+$  ions
- (c) Soluble complex of copper with AgNO3
- (d)  $Cu^-$  ion by the reduction of Cu
- Consider the reaction  $M_{(aq)}^{n+} + ne^- \rightarrow M_{(s)}$ . The standard 14. reduction potential values of the elements  $M_1, M_2$  and  $M_3$  are -0.34V, -3.05V and -1.66V respectively. The order of their reducing power will be [NCERT 1990]
  - (a)  $M_1 > M_2 > M_3$
- (b)  $M_3 > M_2 > M_1$
- (c)  $M_1 > M_3 > M_2$  (d)  $M_2 > M_3 > M_1$
- $E^0 = \frac{RT}{nF}$  In  $K_{eq}$ . This is called 15.

[CPMT 1988; MP PET 2000]

- (a) Gibb's equation
- (b) Gibb's-Helmholtz equation
- (c) Nernst's equation
- (d) Vander Waal's equation
- Four alkali metals A, B, C and D are having respectively standard 16 electrode potential as -3.05,-1.66,-0.40 and 0.80. Which one will be the most reactive

[MP PMT/PET 1988; CPMT 1983: MNR 1993; UPSEAT 2002]

(a) A

(b) B

(c) C

- (d) D
- 17. Which one of the following metals cannot evolve  $H_2$  from acids or  $H_2O$  or from its compounds

[MP PET/PMT 1988; CPMT 1996; AFMC 1998, 99; Pb. PET 1999; BVP 2003]

- (a) Hg
- (b) *Al*
- (c) *Pb*
- (d) Fe
- 18. Which one of the following reaction is not possible

[MP PMT 1991]

- (a)  $Fe + H_2SO_4 \rightarrow FeSO_4 + H_2$
- (b)  $Cu + 2AgNO_3 \rightarrow Cu(NO_3)_2 + 2Ag$
- (c)  $2KBr + I_2 \rightarrow 2KI + Br_2$
- (d)  $CuO + H_2 \rightarrow Cu + H_2O$
- When a rod of metal A is dipped in an aqueous solution of metal 19. B (concentration of  $B^{2+}$  ion being 1M) at  $25^{\circ}$  C, the standard electrode potentials are  $A^{2+}/A = -0.76$  volts,  $B^{2+}/B = +0.34$  volts [KCET 1992]
  - (a) A will gradually dissolve
  - (b) B will deposit on A
  - (c) No reaction will occur
  - (d) Water will decompose into  $H_2$  and  $O_2$

- The reaction  $Zn^{2+} + 2e^{-} \rightarrow Zn$  has a standard potential of 20. -0.76 V. This means [KCET 1992]
  - (a) Zn can't replace hydrogen from acids
  - (b) Zn is a reducing agent
  - (c) Zn is a oxidising agent
  - (d)  $Zn^{2+}$  is a reducing agent
  - $2H^{+}(aq) + 2e^{-} \rightarrow H_{2}(g)$ . The standard electrode potential for the above reaction is (in volts)
    - (a) 0

(b) + 1

(c) - 1

- (d) None of these
- 22. K, Ca and Li metals may be arranged in the decreasing order of their standard electrode potentials as [CPMT 1990]
  - (a) *K*, *Ca*, *Li*
- (b) *Ca*, *K*, *Li*
- (c) Li, Ca, K
- (d) Ca, Li, K
- The correct order of chemical reactivity with water according to electrochemical series [MP PMT 1991]
  - (a) K > Mg > Zn > Cu
- (b) Mg > Zn > Cu > K
- (c) K > Zn > Mg > Cu
- (d) Cu > Zn > Mg > K
- EMF of cell  $Ni \mid Ni^{2+}(1.0 M) \mid Au^{3+}(1.0 M) \mid Au$  (Where 24.  $E^{o}$  for  $Ni^{2+} | Ni$  is  $-0.25 \ V; E^{o}$  for  $Au^{+3} | Au$  is  $1.50 \ V)$  is [MP PET 1993: MP PMT 2000]
  - (a) + 1.25 V
- (b) -1.75 V
- (c) + 1.75 V
- (d) + 4.0 V
- Oxidation and reduction take place in a cell, then its electromotive 25. force will be [RPET 1999]
  - (a) Positive
- (b) Negative
- (c) Zero
- (d) Stable
- For a spontaneous reaction the  $\Delta G$ , equilibrium constant (K) and 26.  $E_{Cell}^{o}$  will be respectively [AIEEE 2005]
  - (a) -ve > 1 + ve
- (b) +ve > 1, -ve
- (c) -ve, < 1, -ve
- (d) -ve, > 1, -ve
- The reference electrode is made from which of the following 27.

[MP PET/PMT 1988]

- (a)  $ZnCl_2$
- (b)  $CuSO_4$
- (c)  $Hg_2Cl_2$
- (d)  $HgCl_2$
- 28. The charge over anode in a galvanic cell is
  - (a) Negative
  - (b) Positive
  - (c) No charge
  - (d) Sometimes negative and sometimes positive
  - The standard electrode potential for the two electrode  $A^+/A$  and  $B^+/B$  are respectively 0.5  $\,V\,$  and 0.75  $\,V\,$  . The  $\,emf\,$  of the given cell  $A \mid A^{+}(a=1) \mid B^{+}(a=1) \mid B$  will be
  - (a) 1.25 V
- (b) -1.25 V
- (c) -0.25 V
- (d) 0.25 V

The standard reduction potential for  $Li^+/Li$ ;  $Zn^{2+}/Zn$ ;

- $H^+/H_2$  and  $Ag^+/Ag$  is -3.05, -0.762, 0.00 and
- +0.80~V. Which of the following has highest reducing capacity
- (a) Ag
- (b)  $H_2$
- (c) Zn
- (d) Li



31. If an iron rod is dipped in  $CuSO_4$  solution

[MADT Bihar 1984]

- (a) Blue colour of the solution turns green
- (b) Brown layer is deposited on iron rod
- (c) No change occurs in the colour of the solution
- (d) Blue colour of the solution vanishes
- (e) None of the above
- **32.**  $E^o$  values of  $Mg^{2+}/Mg$  is -2.37V, of  $Zn^{2+}/Zn$  is -0.76V and  $Fe^{2+}/Fe$  is -0.44V. Which of the following statements is correct [EAMCET 1989]
  - (a) Zn will reduce  $Fe^{2+}$
  - (b) Zn will reduce  $Mg^{2+}$
  - (c) Mg oxidises Fe
  - (d) Zn oxidises Fe
- 33. The standard reduction potential for  $Fe^{2+}/Fe$  and  $Sn^{2+}/Sn$  electrodes are -0.44 and -0.14 volt respectively. For the given cell

reaction  $Fe^{2+} + Sn \rightarrow Fe + Sn^{2+}$ , the standard EMF is [IIT Screening 1990; MP PMT 2011 by salt bridge if

- (a) + 0.30 V
- (b) 0.58 V
- (c) + 0.58 V
- (d) 0.30 V
- **34.** Electrode potential of  $Zn^{2+}/Zn$  is -0.76 V and that of

 ${\it Cu}^{2+}/{\it Cu}$  is  $+0.34{\it V}$  . The  ${\it EMF}$  of the cell constructed between these two electrodes is

[EAMCET 1992; BHU 2001; CBSE PMT 2001; KCET 1990; MHCET 1999, 2003; Pb. CET 2002; AFMC 2001; Pb. PMT 2004]

- (a) 1.10 V
- (b) 0.42 V
- (c) -1.1V
- (d) 0.42 V
- **35.** *EMF* of a cell whose half cells are given below is

$$Mg^{2+} + 2e^{-} \rightarrow Mg(s); E = -2.37 V$$

$$Cu^{2+} + 2e^{-} \rightarrow Cu(s); E = +0.33 V$$

[EAMCET 1987; MP PET 1994; Pb. PMT 2000]

- (a) -2.03 V
- (b) 1.36 V
- (a) -2.03(c) 2.7 V
- (d) 2.03 V
- **36.** A cell constructed by coupling a standard copper electrode and a standard magnesium electrode has *emf* of 2.7 *volts*. If the standard reduction potential of copper electrode is + 0.34 *volt* that of magnesium electrode is [KCET 1989]
  - (a) + 3.04 *volts*
- (b) 3.04 *volts*
- (c) + 2.36 *volts*
- (d) 2.36 *volts*
- 37. When  $E_{Ag^+/Ag}^o = 0.8$  *volt* and  $E_{Zn^{2+}/Zn}^o = -0.76$  *volt*, which of the following is correct [MP PMT 1994]
  - (a)  $Ag^+$  can be reduced by  $H_2$
  - (b) Ag can oxidise  $H_2$  into  $H^+$
  - (c)  $Zn^{2+}$  can be reduced by  $H_2$
  - (d) Ag can reduce  $Zn^{2+}$  ion
- **38.** Adding powdered lead and iron to a solution that is 1.0 M in both  $Pb^{2+}$  and  $Fe^{2+}$  ions, would result a reaction, in which

[CPMT 1987]

- (a) More iron and  $Pb^{2+}$  ions are formed
- (b) More lead and  $Fe^{2+}$  ions are formed
- (c) Concentration of both  ${\it Pb}^{2+}$  and  ${\it Fe}^{2+}$  ions increases
- (d) There is no net change
- **39.** Given standard electrode potentials

$$Fe^{++} + 2e^{-} \rightarrow Fe$$
;  $E^{o} = -0.440 \text{ V}$ 

$$Fe^{+++} + 3e^{-} \rightarrow Fe$$
;  $E^{o} = -0.036 \text{ V}$ 

The standard electrode potential  $(E^o)$  for

$$Fe^{+++} + e^{-} \rightarrow Fe^{++}$$
 is

[AllMS 1982]

- (a) 0.476 V
- (b) 0.404 V
- (c) + 0.404 V
- (d) + 0.771 V
- Reduction potential of four elements P, Q, R, S is -2.90, +0.34, +1.20 and -0.76. Reactivity decreases in the order [MP PET 1989; UPSEAT 2001]
- (a) P > Q > R > S
- (b) Q > P > R > S
- (c) R > Q > S > P
- (d) P > S > Q > R
- **41.** Which of the following metal can deposit copper from copper sulphate solution [CPMT 1983; MP PMT 1989]
  - (a) Mercury
- (b) Iron
- (c) Gold
- (d) Platinum

Standard electrode potential of  $Ag^+/Ag$  and  $Cu^+/Cu$  is +0.80V and +0.34V respectively. these electrodes are joint  $PMT_2003$  by salt bridge if [AMU 2002]

- (a) Copper electrode is work like cathode, then  $E_{cell}^o$  is  $\pm 0.45 V$
- (b) Silver electrode is work like anode then  $E_{cell}^o$  is -0.34V
- (c) Copper electrode is work like anode then  $E_{cell}^o$  is +0.46V
- (d) Silver electrode is work like cathode then  $E_{\it cell}^{\it o}$  is -0.34V
- (e) Silver electrode is work like anode then  $E_{\it cell}^{\it o}$  will be  $+1.14\,V$
- 43. The reaction is spontaneous if the cell potential is

[MP PET 1999]

- (a) Positive
- (b) Negative
- (c) Zero
- (d) Infinite
- **44.** Which substance eliminates bromine from KBr solution

[11T 1981]

(a)  $I_2$ 

- (b) *Cl*<sub>2</sub>
- (c) HI
- (d)  $SO_2$
- **45.** A standard hydrogen electrode has zero electrode potential because[IIT 1997]
  - (a) Hydrogen is easiest to oxidise
  - (b) The electrode potential is assumed to be zero
  - (c) Hydrogen atom has only one electron
  - (d) Hydrogen is the lightest element
- **46.** In the electrochemical cell

$$H_2(g)1 atm |H^+(1M)|| Cu^{2+}(1M)| Cu(s)$$

Which one of the following statements is true

[EAMCET 1997]

- (a)  $H_2$  is cathode; Cu is anode
- (b) Oxidation occurs at Cu electrode
- (c) Reduction occurs at  $H_2$  electrode
- (d)  $H_2$  is anode; Cu is cathode
- **7.** Expression representing the cell potential ( $E \ cell$ )
  - (a)  $E_{a} + E_{a}$
- (b)  $E_{-} E_{-}$
- (c)  $E_{\text{\tiny orbit}} E_{\text{\tiny orbit}}$
- (d)  $E_{\omega} E_{\omega}$
- 48. Iron displaces copper from its salt solution, because

[MP PMT 1996]

(a) Atomic number of iron is less than that of copper



- (b) The standard reduction potential of iron is less than that of copper
- (c) The standard reduction potential of iron is more than that of copper
- (d) The iron salt is more soluble in water than the copper salt
- **49.** (i) Copper metal dissolves in 1*M* silver nitrate solution and crystals of silver metal get deposited.
  - (ii) Silver metal does not react with 1 M zinc nitrate solution
  - (iii) Zinc metal dissolves in 1*M* copper sulphate solution and copper metal gets deposited

Hence the order of decreasing strength of the three metals as reducing agents will be

- (a) Cu > Ag > Zn
- (b) Ag > Cu > Zn
- (c) Zn > Cu > Ag
- (d) Cu > Zn > Ag
- **50.** Standard electrode potentials of Zn and Fe are known to be (i)  $-0.76\,V$  and (ii)  $-0.44\,V$  respectively. How does it explain that galvanization prevents rusting of iron while zinc slowly dissolves away
  - (a) Since (i) is less than (ii), zinc becomes the cathode and iron the anode  $% \left( \frac{1}{2}\right) =\frac{1}{2}\left( \frac{1}{2}\right)$
  - (b) Since (i) is less than (ii), zinc becomes the anode and iron the cathode
  - (c) Since (i) is more than (ii), zinc becomes the anode and iron the cathode
  - (d) Since (i) is more than (ii), zinc becomes the cathode and iron the anode
- Amongst the following electrodes the one with zero electrode potential is [MP PMT 1997]
  - (a) Calomel electrode
  - (b) Standard hydrogen electrode
  - (c) Glass electrode
  - (d) Gas electrode
- **52.** Which of the following is correct expression for electrode potential of a cell [MP PMT 1997]

(a) 
$$E = E^o - \frac{RT}{nF} \ln \frac{[product]}{[reactant]}$$

(b) 
$$E = E^o + \frac{RT}{F} \ln \frac{[product]}{[reactant]}$$

(c) 
$$E = E^o - \frac{RT}{nF} \ln \frac{[\text{reactant}]}{[\text{product}]}$$

(d) 
$$E = -\frac{RT}{F} \ln \frac{[product]}{[reactant]}$$

**53.** Calculate standard free energy change for the reaction  $\frac{1}{2}Cu(s) + \frac{1}{2}Cl_2(g) \Rightarrow \frac{1}{2}Cu^{2+} + Cl^- \text{ taking place at } 25^{\circ}C \text{ in a}$ 

cell whose standard e.m.f. is 1.02 volts [MP PMT 1997]

- (a) -98430 J
- (b) 98430 *J*
- (c) 96500 J
- (d) -49215 J
- 54. In which cell the free energy of a chemical reaction is directly converted into electricity? [MP PET/PMT 1998]
  - (a) Leclanche cell
- (b) Concentration cell
- (c) Fuel cell
- (d) Lead storage battery
- **55.** Nernst equation is related with
  - (a) The electrode potential and concentration of ions in the solution
  - (b) Equilibrium constant and concentration of ions
  - (c) Free energy change and E.M.F. of the cell
  - (d) None of these

**56.** The standard reduction potentials of 4 elements are given below. Which of the following will be the most suitable reducing agent

 $1 = -3.04 \ V$ ,  $11 = -1.90 \ V$ ,  $111 = 0 \ V$ ,  $1V = 1.90 \ V$ 

[CPMT 1999]

(a) 1

(b) 11

(c) 111

- (d) 1V
- **57.** Electrode potential data are given below :

$$Fe^{3+}(aq) + e^{-} \rightarrow Fe^{-1}(aq); E^{o} = +0.77V$$

$$Al^{3+}(aq) + 3e^{-} \rightarrow Al(s); E^{o} = -1.66V$$

$$Br_2(aq) + 2e^- \rightarrow 2Br^-(aq); E^o = +1.08 V$$

Based on the data given above, reducing power of  $Fe^{2+},Al$  and

- $Br^-$  will increase in the order (a)  $Br^- < Fe^{2+} < Al$
- (b)  $Fe^{2+} < Al < Br^{-}$
- (c)  $Al < Br^- < Fe^{2+}$
- (d)  $Al < Fe^{2+} < Br^{-}$
- **58.** The standard electrode potential  $(E^o)$  for  $OCl^-/Cl^-$  and  $Cl^-/\frac{1}{2}Cl_2$  respectively are 0.94~V and -1.36~V. The  $E^o$

value for  $OCl^-/\frac{1}{2}Cl_2$  will be

[KCET 1996]

[Pb. PMT 1998]

- (a) -0.42 V
- (b) -2.20 V
- (c) 0.52 V
- (d) 1.04 V
- **59.** If the reduction potential is more, then

[CPMT 1996]

- (a) It is easily oxidised
- (b) It is easily reduced
- (c) It acts as oxidising agent
- (d) It has redox nature
- **60.** One of the following is false for Hg [BHU 1998]
  - (a) It can evolve hydrogen from  $H_2S$
  - (b) It is a metal
  - (c) It has high specific heat
  - (d) It is less reactive than hydrogen
- **61.**  $E^o$  for the cell  $Zn|Zn^{2+}(aq)||Cu^{2+}(aq)||Cu$  is  $1.10\,V$  at  $25^o\,C$ , the equilibrium constant for the reaction  $Zn + Cu^{2+}(aq) = Cu + Zn^{2+}(aq)$  is of the order of

[CBSE PMT 1997]

- (a)  $10^{-28}$
- (b)  $10^{-37}$
- (c) 10<sup>+18</sup>
- (d) 10<sup>+17</sup>
- **62.** Standard reduction potentials at  $25^{\circ} C$  of  $Li^{+} \mid Li, Ba^{2+} \mid Ba, Na^{+} \mid Na$  and  $Mg^{2+} \mid Mg$  are -3.05, -2.90, -2.71 and -2.37 *volt* respectively. Which one of the following is the strongest oxidising agent

[CBSE PMT 1994; JIPMER 2002]

- (a)  $Na^+$
- (b) *Li*<sup>+</sup>
- (c)  $Ba^{2+}$
- (d)  $Mg^{2+}$
- **63.** Which of the following displaces  $Br_2$  from an aqueous solution containing bromide ions

[CBSE PMT 1994; JIPMER (Med.) 2002]

- (a)  $Cl_2$
- (b) *Cl*
- (c)  $I_2$

- (d)  $I_3^-$
- **64.** For the cell reaction

$$Cu^{2+}(C_1aq) + Zn(s) = Zn^{2+}(C_2aq) + Cu(s)$$



of an	electrochemical	cell,	the	change	in	free	energy	at	a	given
tempe	rature is a functi	on of		[0	CBS	E PM1	Г 1998]			

- (a)  $\ln (C_1)$
- (b)  $\ln (C_2)$
- (c)  $\ln (C_1 + C_2)$
- (d)  $\ln (C_2/C_1)$
- **65.** The e.m.f. of the cell in which the following reaction  $Zn(s) + Ni^{2+}(a=1.0) \Rightarrow Zn^{2+}(a=10) + Ni(s)$  occurs, is found to be 0.5105 V at 298 K. The standard e.m.f. of the cell is

#### [Roorkee Qualifying 1998]

- (a) 0.5400
- (b) 0.4810 V
- (c) 0.5696 V
- (d) -0.5105 V
- **66.** For the redox reaction

 $Zn(s) + Cu^{2+}(0.1M) \rightarrow Zn^{2+}(1M) + Cu(s)$  taking place in a cell,

$$E_{cell}^{o}$$
 is 1.10 *volt.*  $E_{cell}$  for the cell will be  $\left(2.303\frac{RT}{F} = 0.0591\right)$ 

- [AIEEE 2003]
- (a) 2.14 *volt*
- (b) 1.80 volt
- (c) 1.07 volt
- (d) 0.82 volt
- **67.** The *emf* of a Daniel cell at 298K is  $E_1$   $Zn \mid ZnSO_4 \mid \mid CuSO_4 \mid \mid Cu$  when the concentration of  $ZnSO_4$

is 1.0  $\it M$  and that of  $\it CuSO_4$  is 0.01  $\it M$  the  $\it emf$  changed to  $\it E_2$  . What is the relationship between  $\it E_1$  and  $\it E_2$ 

[CBSE PMT 2003]

- (a)  $E_2 = 0 \neq E_1$
- (b)  $E_1 > E_2$
- (c)  $E_1 < E_2$
- (d)  $E_1 = E_2$
- **68.** The oxidation potentials of following half-cell reactions are given  $Zn \rightarrow Zn^{2+} + 2e^{-}$ ;  $E^{o} = 0.76 \text{ V}$ ,

 $Fe \rightarrow Fe^{2+} + 2e^{-}; E^{o} = 0.44 \ V$  what will be the *emf* of cell,

whose cell-reaction is  $Fe^{2+}(aq) + Zn \rightarrow Zn^{2+}(aq) + Fe$ 

[MP PMT 2003]

- (a) -1.20 V
- (b) + 0.32 V
- (c) -0.32 V
- (d) + 1.20 V
- **69.** The  $E^o$  for half cells  $Fe/Fe^{2+}$  and  $Cu/Cu^{2+}$  are 0.44 V and + 0.32 V respectively. Then [MP PMT 2003]
  - (a)  $Cu^{2+}$  oxidises Fe
- (b)  $Cu^{2+}$  oxidises  $Fe^{2+}$
- (c) Cu oxidises  $Fe^{2+}$
- (d) Cu reduces  $Fe^{2+}$
- **70.** What is  $E^o$  for electrode represented by  $Pt, O_2(1 \ atm)/2H^+(Im)$  [JIPMER 1997]
  - (a) Unpredictable
- (b) Zero
- (c) 0.018 V
- (d) 0.118 V
- **71.** The cell potential of a cell in operation is
  - (a) Zero (b)
  - (a) **E**C.0
- (b) Positive
- (c) Negative
- (d) None of the above
- **72.** Which of the following is displaced by Fe [Roorkee 1995]
  - (a) Ag
- (b) *Hg*
- (c) Zn
- (d) Na
- **73.** The standard electrode potential of the half cells are given below  $Zn^{2+} + 2e^- \rightarrow Zn; E = -7.62 V$ ,

$$Fe^{2+} + 2e^{-} \rightarrow Fe; E = -7.81 \text{ V}$$

The *emf* of the cell  $Fe^{2+} + Zn \rightarrow Zn^{2+} + Fe$  is

[CPMT 2003]

- (a) 1.54 V
- (b) -1.54 V

- (c) 0.19 V (d) + 0.19 V
- 74.  $Zn^{2+} + 2e^{-} \rightarrow Zn(s); E^{o} = -0.76$

$$Fe^{3+} + e^{-} \rightarrow Fe^{2+}; E^{o} = -0.77$$

$$Cr^{3+} + 3e^{-} \rightarrow Cr; E^{o} = -0.79$$
,

$$H^+ + 2e^- \rightarrow 1/2H_2$$
;  $E^o = 0.00$ 

Strongest reducing agent is

[BHU 2003]

- (a)  $Fe^{2+}$
- (b) *Zn*
- (c) Cr
- (d)  $H_2$
- **75.** Standard reduction electrode potentials of three metals *A*, *B* and *C* are respectively + 0.5 *V*, 3.0 *V* and 1.2 *V*. The reducing powers of these metals are [IIT 1998; AIEEE 2003]
  - (a) B > C > A
- (b) A > B > C
- (c) C > B > A
- (d) A > C > B
- **76.** For a cell reaction involving a two-electron change, the standard *emf* of the cell is found to be 0.295 V at  $25^{\circ}C$ . The equilibrium constant of the reaction at  $25^{\circ}C$  will be

#### [Roorkee 1999; AIEEE 2003; CBSE PMT 2004]

- (a)  $1\times10^{-10}$
- (b)  $29.5 \times 10^{-2}$

(c) 10

- (d)  $1 \times 10^{10}$
- 77. For the electrochemical cell,
  - $M \mid M^+ \mid X^- \mid X$ ,  $E^o(M^+ / M) = 0.44$  V and  $E^o(X / X^-) = 0.33$  V. From this data one can deduce that

[IIT-JEE (Screening) 2000]

- (a)  $M+X \rightarrow M^+ + X^-$  is the spontaneous reaction
- (b)  $M^+ + X^- \rightarrow M + X$  is the spontaneous reaction
- (c)  $E_{cell} = 0.77 V$
- (d)  $E_{cell} = -0.77 \ V$
- **78.** The standard potential at  $25^{\circ}C$  for the following half reactions are given against them

$$Zn^{2+} + 2e \rightarrow Zn, E^{o} = -0.762V$$

$$Mg^{2+} + 2e \rightarrow Mg, E^{o} = -2.37V$$

When zinc dust is added to the solution of  $MgCl_2$ 

[UPSEAT 2001]

- (a)  $ZnCl_2$  is formed
- (b) Zinc dissolves in the solution
- (c) No reaction takes place
- (d) Mg is precipitated
- **79.**  $KMnO_4$  acts as an oxidising agent in the neutral medium and gets reduced to  $MnO_2$ . The equivalent weight of  $KMnO_4$  in neutral medium [AMU 2001]
  - (a) *mol. wt*/2
- (b) mol.wt/3
- (c) mol. wt/4
- (d) mol.wt/7
- **80.** Which of the following condition will increase the voltage of the cell, represented by the equation

$$Cu_{(s)} + 2Ag^{+}_{(aq)} \rightarrow Cu^{2+}_{(aq)} + 2Ag_{(s)}$$

[CBSE PMT 2001]

- (a) Increase in the concentration of  $Ag^+$ ion
- (b) Increase in the concentration of  $\operatorname{\it Cu}^+$  ion
- (c) Increase in the dimension of silver electrode
- $\begin{tabular}{ll} (d) & \mbox{Increase in the dimension of copper electrode} \\ {\bf 81.} & \mbox{Which will increase the voltage of the cell} \\ \end{tabular}$

$$Sn_{(s)} + 2Ag^{+}_{(aq)} \rightarrow Sn^{2+}_{(aq)} + 2Ag_{(s)}$$

[DPMT 2001]



- (a) Increase in the concentration of  $Ag^+$  ions
- (b) Increase in the concentration of  $Sn^{2+}$  ions
- (c) Increase in size of the silver rod
- 82 The mass of the proton is 1840 times that of electron, its potential difference is V. The kinetic energy of proton is

[DCE 2001]

- (a) 1840 KeV
- (b) 1 *KeV*
- (c)  $\frac{1}{1840}$  KeV
- (d) 920 KeV
- 83. will be given cell  $Pt | H_2(P_1) | H^+_{(aq)} || H_2(P_2) | Pt$ [AIEEE 2002]
  - $\text{(a)} \quad \frac{RT}{f} \log \frac{P_1}{P_2} \qquad \qquad \text{(b)} \quad \frac{RT}{2f} \log \frac{P_1}{P_2}$
  - (c)  $\frac{RT}{f} \log \frac{P_2}{P_2}$  (d) None of these
- What is the potential of a cell containing two hydrogen electrodes 84. the negative one in contact with  $10^{-8} M H^+$  and positive one in contact with 0.025 M H

[MP PMT 2000]

- (a) 0.18 V
- (b) 0.28 V
- (c) 0.38 V
- (d) 0.48 V
- Will  $Fe_{(s)}$  be oxidised to  $Fe^{2+}$  by the reaction with 1 M HCl  $(E^o$ 85.

for  $Fe/Fe^{2+} = +0.44 \text{ V}$ 

[Pb. PMT 2000]

- (a) Yes
- (b) No
- (c) May be
- (d) Can't say
- EMF of a cell in terms of reduction potential of its left and right 86. electrodes is [AIEEE 2002]
  - (a)  $E = E_{left} E_{right}$
- (b)  $E = E_{left} + E_{right}$
- (c)  $E = E_{right} E_{left}$
- (d)  $E = -(E_{right} + E_{left})$
- Arrange the following in the order of their decreasing electrode 87. potential Mg, K, Ba, Ca
  - (a) K, Ba, Ca, Mg
- (b) Ca, Mg, K, Ba
- (c) Ba, Ca, K, Mg
- (d) Mg, Ca, Ba, K
- 88. Which of the following has highest electrode potential

[Pb. PMT 2000]

(a) Li

- (b) *Cu*
- (c) Au
- (d) Al
- 89. The cell reaction of a cell is

$$Mg_{(s)} + Cu^{2+}(aq) \rightarrow Cu_{(s)} + Mg^{2+}(aq)$$

If the standard reduction potentials of Mg and Cu are -2.37and +0.34 V respectively. The *emf* of the cell is

[EAMCET 1995; JIPMER (Med.) 2001;

AFMC 2002: CBSE PMT 20021

- (a) 2.03 V
- (b) -2.03 V
- (c) + 2.71 V
- (d) -2.71 V
- The element which can displace three other halogens from their 90. compound is [EAMCET 1998]
  - (a) Cl
- (b) F
- (c) Br
- (d) I

- Which of the following has been universally accepted as a reference 91. electrode at all temperatures and has been assigned a value of zero volt [AIIMS 1998]
  - (a) Graphite electrode
  - (b) Copper electrode
  - (c) Platinum electrode
  - Standard hydrogen electrode
- Aluminium is more reactive than Fe . But Al is less easily corroded [DCE 1999] than iron because
  - (a) Al is noble metal
  - Iron forms both mono and divalent ions
  - Oxygen forms a protective oxide layer
  - (d) Fe undergoes reaction easily with  $H_2O$
- 93. Zinc displaces copper from the solution of its salt because

[MP PET 1995]

- (a) Atomic number of zinc is more than that of copper
- (b) Zinc salt is more soluble in water than the copper salt
- (c) Gibbs free energy of zinc is less than that of copper
- (d) Zinc is placed higher than copper in electro-chemical series
- 94. An electrochemical cell is set up as follows

 $Pt(H_2, 1 atm)/0.1 M HCl$ 

 $\parallel$  0.1 M acetic acid /(  $H_2, 1 atm$ ) Pt

E.M.F. of this cell will not be zero because

[CBSE PMT 1995]

- (a) The pH of 0.1 M HCl and 0.1 M acetic acid is not the same
- (b) Acids used in two compartments are different
- (c) E.M.F. of a cell depends on the molarities of acids used
- (d) The temperature is constant
- Cu+ ion is not stable in aqueous solution because of 95. disproportionation reaction.  $E^{o}$  value for disproportionation of

(Given 
$$E^o_{Cu^{2+}/Cu^+} = 0.15$$
,  $E^o_{Cu^{2+}/Cu} = 0.34V$ ) [IIT 1998]

- (a) -0.49 V
- (b) 0.49 V
- (c) -0.38 V
- (d) 0.38 V
- $E^o$  of a cell  $aA + bB \rightarrow cC + dD$  is 96.

(a) 
$$-\frac{RT}{nF}\log\frac{[C]^c[D]^d}{[A]^a[B]^b}$$
 (b)  $-RT\log\frac{[a]^A[b]^B}{[a]^C[d]^D}$ 

(c) 
$$-\frac{RT}{nF}\log\frac{\left[C\right]^{c}\left[d\right]^{D}}{\left[A\right]^{a}\left[B\right]^{b}}$$
 (d)  $-\frac{RT}{nF}\log\frac{\left[C\right]^{c}\left[d\right]^{D}}{\left[a\right]^{A}\left[B\right]^{b}}$ 

In the experiment set up for the measurement of EMF of a half cell 97. using a reference electrode and a salt bridge, when the salt bridge is removed, the voltage

[NCERT 1984; CPMT 1985]

- (a) Does not change
- (b) Decreases to half the value
- Increase to maximum
- (d) Drops to zero
- 98. Electrode potentials of five elements A, B, C, D and E are respectively - 1.36, - 0.32, 0, - 1.26 and -0.42. The reactivity order of these elements are in the order of

[MP PMT 1995]

- (a) A, D, E, B and C
- (b) *C, B, E, D* and *A*
- (c) B, D, E, A and C
- (d) C, A, E, D and B
- What is wrongly stated about electrochemical series 99.

## UNIVERSAL SELF SCORER

#### **522 Electrochemistry**

[DCE 1999]

- (a) It is the representation of element in order of increasing or decreasing standard electrode reductional potential
- (b) It does not compare the relative reactivity of metals
- (c) It compares relative strengths of oxidising agents
- (d)  $H_2$  is centrally placed element

100. Which of the following statements is true for fuel cells

rue for fuel cells
[KCET (Med.) 1999; AFMC 2000]
110. Standar

- (a) They are more efficient
- (b) They are free from pollution
- (c) They run till reactants are active
- (d) All of these

101. What is the potential of a half-cell consisting of zinc electrode in 0.01m  $ZnSO_4$  solution at  $258^oC$  ( $E^o=0.763V$ )

[AIIMS 2000; BHU 2000]

- (a) 0.8221 V
- (b) 8.221 V
- (c) 0.5282 V
- (d) 9.232 V

**102.** The *emf* of a galvanic cell, with electrode potentials of silver = +0.80 V and that of copper = +0.34 V, is **[AIIMS 1999]** 

- (a) -1.1 V
- (b) + 1.1 V
- (c) + 0.46 V
- (d) + 0.76 V

103. Copper cannot replace..... from solution [DPMT 2002]

(a)  $F\epsilon$ 

- (b) *Au*
- (c) Hg
- (d) Ag

104. The strongest reducing agent of the alkali metal is

[CBSE PMT 2000]

(a) Li

(b) *Na* 

(c) K

(d) *Cs* 

105. Which of the following is the most electropositive element

[Pb. PMT 2000]

- (a) Carbon
- (b) Calcium
- (c) Chlorine
- (d) Potassium

**106.** The metal that forms a self protecting film of oxide to prevent corrosion, is [BHU 1999]

- (a) *Cu*
- (b) *Al*
- (c) Na
- (d) Au

107. In a cell that utilises the reaction  $Zn_{(s)} + 2H^{+}(aq) \rightarrow$ 

 $Zn^{2+}(aq) + H_{2(g)}$  addition of  $H_2SO_4$  to cathode compartment, will

- (a) Increase the E and shift equilibrium to the right
- (b) Lower the *E* and shift equilibrium to the right
- (c) Lower the E and shift equilibrium to the left
- (d) Increase the E and shift equilibrium to the left

- (a)  $E^{\circ}_{cell} = -0.77 V$
- (b)  $M^+ + X^- \rightarrow M + X$  is the spontaneous reaction
- (c)  $M + X \rightarrow M^+ + X^-$  is the spontaneous reaction
- (d)  $E^{\circ}_{cell} = .77 V$

**109.** The standard *e.m.f.* of a call, involving one electron change is found to be 0.591 V at  $25^{\circ}C$ . The equilibrium constant of the reaction is

 $(F = 96,500 \ C \ mol^{-1}; \ R = 8.314 \ JK^{-1}mol^{-1})$  [AIEEE 2004]

- (a)  $1.0 \times 10^{10}$
- (b)  $1.0 \times 10^5$
- (c)  $1.0 \times 10^{1}$
- (d)  $1.0 \times 10^{30}$

110. Standard electrode potential of cell  $\ H_2 \ | \ H^+ \ | \ Ag^+ \ | \ Ag$  is

[AIEEE 2004]

- (a) 0.8 V
- (b) -0.8 V
- (c) -1.2 V
- (d) 1.2 V

III. A galvanic cell with electrode potential of ' $A'=+2.23\ V$  and ' $B'=-1.43\ V$ '. The value of  $E^{\circ}_{cell}$  is [Pb.CET 2003]

- (a) 3.66 V
- (b) 0.80 V
- (c) -0.80 V
- (d) -3.66 V

112. The *e.m.f.* of a cell whose half cells are given below is  $Mg^{2+} + 2e^{-} \rightarrow Mg(s) E^{\circ} = -2.37 V$ 

$$Cu^{2+} + 2e^{-} \rightarrow Cu(s) E^{\circ} = +0.34 V$$

[Pb.CET 2001]

- (a) + 1.36 V
- (b) + 2.71 V
- (c) + 2.17 V
- (d) -3.01 V

113. For the cell reaction,  $2Ce^{4+} + Co \rightarrow 2Ce^{3+} + Co^{2+}$   $E^{\circ}_{cell}$  is 1.89 *V*. If  $E^{\circ}_{Ce^{4+}/Ce^{3+}}$  [Pb.CET 2000]

- (a) -1.64 V
- (b) + 1.64 V
- (c) -2.08 V
- (d) + 2.17 V

114. If the  $\Delta G$  of a cell reaction  $AgCl + e^- \rightarrow Ag + Cl^-$  is  $-21.20 \ KJ$ ; the standard *e.m.f.*, of cell is [MP PMT 2004]

- (a) 0.229 V
- (b) 0.220 V
- (c) -0.220 V
- (d) 0.110 V

115. The *e.m.f.* of the cell  $Ag | Ag^+(0.1M)| | Ag^+(1M)| Ag$  at 298 *K* is [DCE 2003]

- (a) 0.0059 V
- (b) 0.059 V
- (c) 5.9 V
- (d) 0.59 V

**116.** The *e.m.f.* of the cell  $Zn|Zn^{2+}(0.01M)||Fe^{2+}(0.001M)||Fe$  at 298 K is 0.2905 then the value of equilibrium for the cell reaction is [IIT-JEE Sci

- (a)  $\frac{0.32}{a^{0.0295}}$
- (b)  $\frac{0.32}{10^{0.0295}}$
- (c)  $\frac{0.26}{10^{0.0295}}$
- (d)  $\frac{0.32}{10^{0.0591}}$

117. Aluminium displaces hydrogen from dilute HCl whereas silver does not. The *e.m.f.* of a cell prepared by combining  $Al/Al^{3+}$  and  $Ag/Ag^+$  is 2.46 V. The reduction potential of silver electrode is +0.80~V. The reduction potential of aluminium electrode is [KCET 2004]

- (a) +1.66 V
- (b) -3.26 V
- (c) 3.26 V
- (d) -1.66 V

118. Consider the following  $E^0$  values:

$$E^0_{Fe^{3+}/Fe^{2+}} = +0.77 \ V$$

$$E^0_{Sn^{2+}/Sn} = -0.14 V$$



Under standard conditions the potential for the reaction  $Sn_{(s)} + 2Fe^{3+}(aq) \rightarrow 2Fe^{2+}(sq) + Sn^{2+}(aq)$  is [AIEEE 2004]

- (a) 0.91 V
- (b) 1.40 V
- (c) 1.68 V
- (d) 0.63 V

119. 
$$Cr_2O_7^{2-} + I^- \rightarrow I_2 + Cr^{3+}$$

$$E^{0}_{cell} = 0.79 V$$

$$E_{Cr_2Q_2^{2-}}^0 = 1.33 \ V, E_{I_2}^0$$
 is

[BVP 2004]

- (a) -0.10 V
- (b) +0.18 V
- (c) -0.54 V
- (d) 0.54 V

120. 
$$Zn(s) + Cl_2(1 \text{ atm}) \rightarrow Zn^{2+} + 2Cl^-$$
.  $E^0_{cell}$  of the cell is 2.12  $V$ . To increase  $E$  [BVP 2004]

- (a)  $[Zn^{2+}]$  should be increased
- (b)  $[Zn^{2+}]$  should be decreased
- (c)  $[Cl^-]$  should be decreased
- (d)  $P_{Cl_2}$  should be decreased

121. The 
$$E^0_{M^{3+}/M^{2+}}$$
 values for  $Cr, Mn, Fe$  and  $Co$  are  $-0.41, +1.57, +0.77$  and  $+1.97$   $V$  respectively. For which one of these metals the change in oxidation state from  $+2$  to  $+3$  is easiest [AIEEE 2004]

- (a) *Fe*
- (b) *Mn*

(c) Cr

(d) *Co* 

122. The rusting of iron takes place as follows

$$2H + 2e^- + \frac{1}{2}O_1 \longrightarrow HO(1)$$
;

$$E^{\circ} = +1.23 \ V$$

$$Fe^{c} + 2e^{-} \longrightarrow Fe(s)$$
;  $E^{b} = -0.44 V$ 

Calculate  $\Delta G$  for the net process

[IIT 2005]

- (a) −322 *kJ mol*
- (b) -161 kJ mol
- (c) -152 kJ mol
- (d) -76 *kJ mol*
- 123. When an acid cell is charged, then

[AFMC 2005]

- (a) Voltage of cell increases
  - (b) Electrolyte of cell dilutes
  - (c) Resistance of cell increases
  - (d) None of these
- 124. The standard electrode potential is measured by

[KCET 2005]

- (a) Electrometer
- (b) Voltmeter
- (c) Pyrometer
- (d) Galvanometer

125. Aluminium displaces hydrogen from acids but copper does not. A galvanic cell prepared by combining 
$$Cu/Cu^{2+}$$
 and  $Al/Al^{3+}$  has an *e.m.f.* of 2.0  $V$  at 298  $K$ . If the potential of copper electrode is + 0.34  $V$ , that of aluminium is

[CPMT 2001; KCET 2001]

- (a) + 1.66 V
- (b) -1.66 V
- (c) + 2.34 V
- (d) -2.3 V

126. If the standard electrode potential of 
$$Cu^{2+}/Cu$$
 electrode is 0.34  $V$ , what is the electrode potential of 0.01 $M$  concentration of

$$Cu^{2+}$$
  $(T = 298 K)$ 

#### [EAMCET 2003]

- (a) 0.399 V
- (b) 0.281 V
- (c) 0.222 V
- (d) 0.176 V

**127.** Calculate the electrode potential at  $298^{\circ}K$  for  $Zn \mid Zn^{++}$  electrode in which the activity of zinc ions is 0.001 M and

 $E^{o}_{Zn/Zn^{++}}$  is -0.74 volts

[AMU 2002]

- (a) 0. 38 volts
- (b) 0.83 *volts*
- (c) 0.40 volts
- (d) 0.45 volts
- 128. Which of the following expression is correct

[Orissa JEE 2005]

- (a)  $\Delta G^o = -nFE^o_{cell}$
- (b)  $\Delta G^o = +nFE^o_{cell}$
- (c)  $\Delta G^o = -2.303RT \, nFE^o_{cell}$  (d)  $\Delta G^o = -nF \log K_C$
- 129. For the feasibility of a redox reaction in a cell, the e.m.f. should be[] & K 2002]
  - (a) Positive
- (b) Fixed
- (c) Zero
- (d) Negative

#### Corrosion

. Corrosion is basically a

[Kerala (Med.) 2002]

- (a) Altered reaction in presence of  $\,H_2O\,$
- (b) Electrochemical phenomenon
- (c) Interaction
- (d) Union between light metal and heavy metal
- 2. Rusting of iron is catalysed by which of the following

[MNR 1990; UPSEAT 2001]

- (a) Fe
- (b)  $O_2$
- (c) Zn

- 1) H+
- 3. Which of the following is a highly corrosive salt

[AFMC 2005]

- (a)  $FeCl_2$
- (b)  $PbCl_2$
- (c)  $Hg_2Cl_2$
- (d)  $HgCl_2$
- **4.** Corrosion of iron is essentially an electrochemical phenomenon where the cell reactions are **[KCET 2005]** 
  - (a) Fe is oxidised to  $Fe^{2+}$  and dissolved oxygen in water is reduced to OH
  - (b) Fe is oxidised to  $Fe^{3+}$  and  $H_2O$  is reduced to  $O_2^{2-}$
  - (c) Fe is oxidised to  $Fe^{2+}$  and  $H_2O$  is reduced to  $O_2^-$
  - (d) Fe is oxidised to  $Fe^{2+}$  and  $H_2O$  is reduced to  $O_2$

# Critical Thinking

## **Objective Questions**

- The limiting molar conductivities  $\wedge^0$  for *NaCl, KBr* and *KCl* are 126, 152 and 150  $S \ cm^2 mol^{-1}$  respectively. The  $\wedge^0$  for *NaBr* is
  - (a)  $278 S cm^2 mol^{-1}$
- (b)  $176 S cm^2 mol^{-1}$
- (c)  $128 S cm^2 mol^{-1}$
- (d)  $302 S cm^2 mol^{-1}$
- On the basis of the electrochemical theory of aqueous corrosion, the reaction occurring at the cathode is

[MP PET 1994; UPSEAT 2001]

- (a)  $O_{2(g)} + 4H_{(aq)}^+ + 4e^- \rightarrow 2H_2O_{(l)}$
- (b)  $Fe_{(s)} \to Fe_{(aa)}^{2+} + 2e^{-}$
- (c)  $Fe_{(aa)}^{2+} \to Fe_{(aa)}^{3+} + e^{-}$



- (d)  $H_{2(g)} + 2OH_{(aq)}^- \rightarrow 2H_2O_{(l)} + 2e^{-1}$
- 3. The reaction  $\frac{1}{2}H_2(g) + AgCl(s) \rightarrow H^+(aq) + Cl^-(aq) + Ag(s)$  occurs in the galvanic cell

[IIT 1985; AMU 2002; KCET 2003]

- (a)  $Ag/AgCl(s)KCl(soln)||AgNO_3(soln)/Ag$
- (b)  $Pt/H_2(g)HCl(soln)|| AgNO_3(soln)/Ag$
- (c)  $Pt/H_2(g)HCl(soln)|| AgCl(s)/Ag$
- (d)  $Pt/H_2(g)KCl(soln)|| AgCl(s)/Ag$
- **4.** The standard reduction potential  $E^o$  for the half reactions are as

$$Zn = Zn^{2+} + 2e^{-}; E^{o} = +0.76 V$$

$$Fe = Fe^{2+} + 2e^{-}$$
;  $E^{o} = +0.41 V$ 

The *EMF* for cell reaction  $Fe^{2+} + Zn \rightarrow Zn^{2+} + Fe$  is

[IIT 1988; CBSE PMT 1993, 96; BHU 1995, 2000; CPMT 2000; KCET 2000; AllMS 2001; Orissa JEE 2002]

- (a) -0.35 V
- (b) +0.35 V
- (c) +1.17 V
- (d) -1.17 V
- **5.** The number of electrons to balance the following equation  $NO_3^- + 4H^+ + e^- \rightarrow 2H_2O + NO$  is [IIT Screening 1991]
  - (a) 5

(b) 4

(c) 3

- (d) 2
- **6.** The standard EMF for the given cell reaction  $Zn + Cu^{2+} = Cu + Zn^{2+}$  is 1.10V at  $25^{\circ}C$ . The EMF for the cell reaction, when 0.1M  $Cu^{2+}$  and 0.1M  $Zn^{2+}$  solutions are used, at  $25^{\circ}C$  is

[MNR 1994; AMU 1999; UPSEAT 2002]

- (a) 1.10 V
- (b) 0.110 V
- (c) -1.10 V
- (d) -0.110 V
- 7. A gas X at 1 atm is bubbled through a solution containing a mixture of  $1MY^-$  and  $1MZ^-$  at  $25^{\circ}C$ . If the reduction potential of Z>Y>X, then [IIT 1999]
  - (a) Y will oxidize X and not Z
  - (b) Y will oxidize Z and not X
  - (c) Y will oxidize both X and Z
  - (d) Y will reduce both X and Z
- **8.** The oxidation potential of a hydrogen electrode at pH=10 and  $pH_1=1$  [JIPMER 2000]
  - (a) 0.059 V
- (b) 0.59 V
- (c) 0.00 V
- (d) 051 1/
- **9.** The decomposition of hydrogen peroxide is an example of

[Roorkee 2000]

- (a) Exothermic reaction
- (b) Endothermic reaction
- (c) Negative catalysis
- (d) Auto-oxidation
- **10.** Aluminium oxide may be electrolysed at 1000°C to furnish aluminium metal (At. Mass = 27 amu; 1 Faraday = 96,500 Coulombs). The cathode reaction is

$$Al^3 + 3e^- \rightarrow Al^\circ$$

To prepare 5.12kg of aluminium metal by this method would require [AIEEE 2005]

- (a)  $5.49 \times 10^7 C$  of electricity
- (b)  $1.83 \times 10^7 C$  of electricity
- (c)  $5.49 \times 10^4 C$  of electricity
- (d)  $5.49 \times 10^{1} C$  of electricity

Electrolyte :	KCl	KNO <sub>3</sub>	HCl	NaOAc	NaCl
$\Lambda^{\infty}(Scm^{2}mof^{1}):$	149.9	145.0	426.2	91.0	126.5

Calculate  $\Lambda_{HOAc}^{\infty}$  using appropriate molar conductances of the electrolytes listed above at infinite dilution in  $H_2O$  at 25° **C[AIEEE 2005]** 

(a) 517.2

11.

- (b) 552.7
- (c) 390.7
- (d) 217.5
- 12. The mass of carbon anode consumed (giving only carbondioxide) in the production of 270kg of aluminium metal from bauxite by the Hall process is [CBSE PMT 2005]
  - (a) 180kg
- (b) 270 kg
- (c) 540 kg
- (d) 90kg
- 13. 4.5g of aluminium (at mass 27amu) is deposited at cathode from  $Al^{3+}$  solution by a certain quantity of electric charge. The volume of hydrogen produced at STP from  $H^+$  ions in solution by the same quantity of electric charge will be

[CBSE PMT 2005]

- (a) 22.4 L
- (b) 44.8 *L*
- (c) 5.6 L
- (d) 11.2 L
- 14. What amount of *Cl* gas liberated at anode, if 1 *amp*. current is passed for 30 *min*. from *NaCl* solution.

[BHU 2005]

- (a) 0.66 moles
- (b) 0.33 moles
- (c) 0.66 g
- (d) 0.33 g



Read the assertion and reason carefully to mark the correct option out of the options given below:

- (a) If both assertion and reason are true and the reason is the correct explanation of the assertion.
- (b) If both assertion and reason are true but reason is not the correct explanation of the assertion.
- (c) If assertion is true but reason is false.
- (d) If the assertion and reason both are false.
- (e) If assertion is false but reason is true.
- Assertion : Sodium ions are discharged in preference to hydrogen ions at a mercury cathode.

Reason : The nature of the cathode can effect the order of

discharge of ions.

 Assertion : In electrolysis, the quantity of electricity needed for depositing 1 mole of silver is different from

that required for 1 mole of copper.

Reason : The molecular weights of silver and copper are

Assertion : Equivalent conductance of all electrolytes decreases with the increases in concentration.

Reason : Lesser number of ions are available per gram equivalent at higher concentration.

. Assertion : Copper reacts with hydrochloric acid and liberates hydrogen from the solution of dilute

hydrochloric acid. Reason : Hydrogen is below copper in the electrochemical

series.



5.	Assertion	:	Copper liberates hydrogen from a solution of dil.
			HCl.
	Reason		Hydrogen is below copper in the reactivity series

	Reason	:	Hydrogen is below copper in the reactivity series.
6.	Assertion	:	Zn metal is formed when a $Cu$ plate in dipped in

 $ZnSO_4$  solution. Reason : Cu being placed above Zn in electrochemical

7. Assertion : Electrical conductivity of copper increases with increase in temperature.

Reason : The electrical conductivity of metals is due to the motion of electrons.

**8.** Assertion : A small amount of acid or alkali is added before electrolysis of water.

Reason : Pure water is weak electrolyte.

9. Assertion : Copper reacts with HCl and liberates hydrogen.Reason : Hydrogen is present above Cu in the reactivity

10. Assertion : K and Cs are used in photoelectric cells.

Reason : K and Cs emit electrons on exposure to ligh

Reason : K and Cs emit electrons on exposure to light.

11. Assertion : A large dry cell has high e.m.f.

Reason : The *e.m.f.* of a dry cell is proportional to its size.

12. Assertion : The resistivity for a substance is its resistance when its is one meter long and its area of cross section is one square meter.

Reason : The SI uints of resistivity are ohm metre  $(\Omega m)$  and ohm centimeter  $(\Omega cm)$ .

13. Assertion : When acidified zinc sulphate solution is electrolysed between zinc electrodes, it is zinc that is deposited at the cathode and hydrogen evolution does not take place.

Reason : The electrode potential of zinc is more negative than hydrogen as the overvoltage for the hydrogen as the evolution on zinc is quite large.

14. Assertion : If  $\lambda_{Na^+}^0 + \lambda_{Cl^-}^0$  are molar limiting conductivity of the sodium and chloride ions respectively, then the limiting molar conducting for sodium chloride is given by the equation:

 $\wedge_{NaCl}^0 = \lambda_{Na^+}^0 + \lambda_{Cl^-}^0 \, .$ 

Reason : This is according to Kohlrausch law of

independent migration of ions.

15. Assertion : One coulomb of electric charge deposits weight equal to the electrochemical equivalent of the

substance.

16.

18.

21.

Reason

Reason : One Faraday deposits one mole of the substance.

Assertion : Auric chloride (AuCl<sub>3</sub>) solution cannot be

Auric chloride  $(AuCl_3)$  solution cannot be stored in a vessel made of copper, iron, nickel, chromium, zinc or tin.

Reason : Gold is a very precious metal.

17. Assertion : For a cell reaction  $Zn(s) + Cu^{2+}(aq) \rightarrow Zn^{2+}(aq) + Cu(s)$ ; at the equilibrium,

voltmeter gives zero reading. At the equilibrium, there is no change in the

concentration of  $Cu^{2+}$  and  $Zn^{2+}$  ions.

Assertion : A negative value of standard reduction potential means that reduction take place on this electrode

with reference to standard hydrogen electrode.

: The standard electrode potential of a half cell has

Reason : The standard electrode potential of a a fixed value.

Assertion : Weston is a standard cell.

19. Assertion : Weston is a standard cell.Reason : Its e.m.f. does not change with temperature.

**20.** Assertion : Galvanic cells containing hydrogen, methane, methanol etc. as fuels are called fuel cells.

Reason : They are designed to convert the energy of combustion of fuels directly into electrical

energy.

Assertion : Zinc displaces copper from copper sulphate

Reason : The  $E^0$  of zinc is -0.76 V and that of copper

is +0.34 V.

22. Assertion : Identification of cathode and anode is done by

the use of a thermometer.

Reason : Higher is the value of reduction potential, greater

would be its reducing power.

23. Assertion : An electrochemical cell can be set up only if the

redox reaction is spontaneous.

Reason : A reaction is spontaneous if free energy change is

negative.

**24.** Assertion : Galvanised iron does not rust.

Reason : Zinc has a more negative electrode potential than iron. [AIIMS 2005]

**25.** Assertion : In an electrochemical cell anode and cathode are respectively negative and positive electrodes.

respectively negative and positive electrodes.

Reason : At anode oxidation takes place and at cathode

reduction takes place and a

**26.** Assertion :  $Ni/Ni^{2+}(1.0M)||Au^{3+}(1.0M)||Au$ , for this

cell emf is 1.75 V if  $E^o_{Au^{3+}/Au}=1.50$  and

 $E_{Ni^{2+}/Ni}^o = 0.25V$ .

Reason : Emf of the cell =  $E_{\text{cathode}}^o - E_{\text{anode}}^o$ .

**27.** Assertion : Salts like  $KCl, KNO_3$  i.e., inert electrolytes are

used in salt bridge.

Reason : An inert electrolyte can easily be filled in the U-

tube.

**28.** Assertion : Emf and potential difference are same for cell.

Reason : Both gives the difference in electrode potential

under any condition.



## **Electrolytes and Electrolysis**

1	b	2	d	3	С	4	С	5	d
6	d	7	b	8	b	9	b	10	а
11	а	12	С	13	d	14	d	15	С
16	d	17	а	18	С	19	С	20	а
21	а	22	b	23	b	24	d	25	b
26	b	27	b	28	а	29	С	30	b
31	d	32	d	33	С	34	b	35	b
36	а	37	d	38	С	39	d	40	а
41	а								

### Faraday's law of electrolysis

1	С	2	а	3	b	4	С	5	С
6	С	7	С	8	b	9	С	10	а
11	С	12	а	13	С	14	b	15	b

	TIP DO THE
	IVERSAL
SEL	F SCORER

16	С	17	а	18	а	19	d	20	С
21	d	22	С	23	b	24	d	25	а
26	d	27	а	28	d	29	С	30	а
31	b	32	b	33	а	34	b	35	b
36	b	37	b	38	b	39	а	40	а
41	С	42	а	43	а	44	b	45	С
46	b	47	d	48	d	49	b	50	а
51	С	52	b	53	b	54	а	55	С
56	С	57	С	58	d	59	d	60	d
61	С	62	С	63	b	64	а	65	b
66	b	67	а	68	а	69	b	70	С
71	С	72	b	73	а	74	b	75	С
76	b	77	С	78	С	79	b	80	b
81	С	82	b	83	а	84	С	85	С
86	е	87	b	88	а	89	d	90	С
91	С	92	а	93	С	94	b		

Cana	luctor	and	Conductance	
COLIC	luctor	anu	Conductance	;

1	b	2	b	3	d	4	d	5	а
6	b	7	b	8	а	9	b	10	d
11	а	12	b	13	b	14	b	15	b
16	d	17	b	18	d	19	b	20	С
21	b	22	а	23	С	24	d	25	a
26	b	27	d						

## **Cell constant and Electrochemical cells**

	-								
1	d	2	d	3	d	4	b	5	b
6	а	7	С	8	С	9	b	10	b
11	С	12	С	13	а	14	а	15	d
16	С	17	С	18	а	19	b	20	b
21	b	22	b	23	а	24	b	25	С
26	а	27	С	28	а	29	b	30	С
31	а	32	b	33	С	34	b	35	d
36	b	37	С	38	d	39	b	40	d
41	а	42	d	43	С	44	а	45	d
46	а	47	d	48	b	49	С	50	С
51	b	52	d	53	С	54	а	55	С
56	d	57	b	58	а	59	а	60	С
61	а								

## Electrode potential, E<sub>Cell</sub>, Nernst equation and ECS

1	b	2	а	3	а	4	b	5	d
6	С	7	С	8	а	9	а	10	b
11	а	12	С	13	a	14	d	15	С

16	а	17	а	18	С	19	b	20	b
21	а	22	b	23	а	24	С	25	а
26	а	27	С	28	а	29	d	30	d
31	b	32	а	33	d	34	а	35	С
36	d	37	а	38	b	39	d	40	d
41	b	42	С	43	а	44	b	45	b
46	d	47	С	48	b	49	С	50	b
51	b	52	а	53	а	54	С	55	а
56	а	57	а	58	a	59	С	60	a
61	b	62	d	63	а	64	d	65	b
66	С	67	b	68	b	69	а	70	b
71	b	72	ab	73	С	74	С	75	a
76	d	77	b	78	С	79	b	80	а
81	а	82	b	83	b	84	С	85	а
86	С	87	а	88	С	89	С	90	b
91	d	92	С	93	d	94	а	95	d
96	а	97	d	98	a	99	b	100	d
101	а	102	С	103	a	104	а	105	d
106	b	107	а	108	b	109	а	110	a
111	а	112	b	113	b	114	b	115	b
116	b	117	d	118	а	119	d	120	b
121	С	122	а	123	а	124	b	125	b
126	b	127	b	128	а	129	а		

## Corrosion

1	b	2	d	3	d	4	а	

## **Critical Thinking Questions**

1	С	2	а	3	С	4	b	5	С
6	а	7	а	8	b	9	acd	10	а
11	С	12	d	13	С	14	С		

## Assertion & Reason

1	a	2	b	3	а	4	d	5	d
6	d	7	е	8	а	9	е	10	а
11	d	12	b	13	а	14	а	15	С
16	b	17	а	18	е	19	а	20	а
21	a	22	d	23	b	24	а	25	а
26	а	27	С	28	d				