CY : CHEMISTRY

Duration : Three Hours

Maximum Marks :100

CY

Read the following instructions carefully.

- This question paper contains 16 printed pages including pages for rough work. Please check all pages and report discrepancy, if any.
- Write your registration number, your name and name of the examination centre at the specified locations on the right half of the Optical Response Sheet (ORS).
- Using HB pencil, darken the appropriate bubble under each digit of your registration number and the letters corresponding to your paper code.
- 4. All questions in this paper are of objective type.
- 5. Questions must be answered on Optical Response Sheet (ORS) by darkening the appropriate bubble (marked A, B, C, D) using HB pencil against the question number on the left hand side of the ORS. Each question has only one correct answer. In case you wish to change an answer, erase the old answer completely. More than one answer bubbled against a question will be treated as an incorrect response.
- 6. <u>There are a total of 60 questions carrying 100 marks.</u> Questions 1 through 20 are 1-mark questions, questions 21 through 60 are 2-mark questions.
- 7. Questions 51 through 56 (3 pairs) are common data questions and question pairs (57, 58) and (59, 60) are linked answer questions. The answer to the second question of the above 2 pairs depends on the answer to the first question of the pair. If the first question in the linked pair is wrongly answered or is un-attempted, then the answer to the second question in the pair will not be evaluated.
- 8. Un-attempted questions will carry zero marks.
- 9. Wrong answers will carry NEGATIVE marks. For Q.1 to Q.20, ¹/₃ mark will be deducted for each wrong answer. For Q. 21 to Q. 56, ³/₃ mark will be deducted for each wrong answer. The question pairs (Q.57, Q.58), and (Q.59, Q.60) are questions with linked answers. There will be negative marks only for wrong answer to the first question of the linked answer question pair i.e. for Q.57 and Q.59, ³/₃ mark will be deducted for each wrong answer. There is no negative marking for Q.58 and Q.60.
- 10. Calculator (without data connectivity) is allowed in the examination hall.
- 11. Charts, graph sheets or tables are NOT allowed in the examination hall.
- Rough work can be done on the question paper itself. Additionally, blank pages are given at the end of the question paper for rough work.

Some Useful Data

Boltzmann constant, $k_B = 1.381 \times 10^{-23} \text{ J K}^{-1}$

Avogadro number, $N = 6.022 \times 10^{23} \text{ mol}^{-1}$

Electron charge, $e = 1.602 \times 10^{-19} C$

Faraday constant, $F = 96500 \text{ C mol}^{-1}$

Planck constant, $h = 6.626 \times 10^{-34} \text{ J s}$

Speed of light, $c = 2.998 \times 10^8 \text{ m s}^{-1}$

2009					CY		
Q. 1 -	- Q. 20 carry one r	nark each.					
Q.1	The ³¹ P NMR spectr	rum of P ₄ S ₃ consists of					
	(A) a singlet		(B) a doublet and a t	riplet			
	(C) a doublet and a c	quartet	(D) two doublets	In the reaction			
Q.2	The geometry aroun	d the central atom in ClF4	* is				
	(A) square planar	(B) square pyramidal	(C) octahedral	(D) trigonal bipyramidal			
Q.3	The correct statement	nt about the Cu-N bond di	stances in [Cu(NH ₃) ₆]	²⁺ is			
	(A) all the bond dist	ances are equal					
	(B) the axial bonds a	are longer than the equator	rial ones				
	(C) the equatorial be	onds are longer than the ax	cial ones				
	(D) all the bond dist	ances are unequal					
Q.4	The reaction of phosgene with an excess of NH ₃ produces						
	(A) HN=C=O	(B) H ₂ N–C(Cl)=O	(C) (H ₂ N) ₂ C=O	(D) (H ₂ N) ₂ CCl ₂			
Q.5	The number of meta	al – metal bonds in $[(C_5H_5)]$)Fe(CO)] ₂ is				
	(A) zero	(B) one	(C) two	(D) three			
Q.6	The coordination number of the Ba^{2+} ions in barium fluoride is 8. The coordination number of the fluoride ion is						
	(A) 8	(B) 4	(C) 1	(D) 2			
Q.7	In the transformation of oxyhaemoglobin to deoxyhaemoglobin						
	(A) Fe ²⁺ in the low spin state changes to Fe ²⁺ in the high spin state						
	(B) Fe ²⁺ in the low spin state changes to Fe ³⁺ in the low spin state						
	(C) Fe ²⁺ in the high spin state changes to Fe ²⁺ in the low spin state						
	(D) Fe^{2+} in the high	spin state changes to Fe ³⁺	in the high spin state				
Q.8	For the compound						
	ОН						
	Ö the stereochemical	notations are					
	(A) 2Z, 4R	(B) 2Z, 4S	(C) 2E, 4R	(D) 2 <i>E</i> , 4 <i>S</i>			
CY					3/16		

2009





(A) aromatic and has high dipole moment

(C) non-aromatic and has high dipole moment

excess

[X]

- (B) aromatic and has no dipole moment
- (D) anti-aromatic and has no dipole moment

In the reaction Q.10

the major product X is





0.11 In the reaction

Δ

 $(* = {}^{13}C$ labelled carbon)

the major products X and Y are



[X] +[Y

Q.12 In the reaction

the major product X is



CY



- Q.19 According to the Debye-Hückel limiting law, if the concentration of a dilute aqueous solution of KCl is increased 4-fold, the value of $\ln \gamma_{\pm} (\gamma_{\pm}$ is the molal mean ionic activity coefficient) will
 - (A) decrease by a factor of 2

- (B) increase by a factor of 2
- (C) decrease by a factor of 4
- (D) increase by a factor of 4
- Q.20 For the parallel first order reaction shown below

$$X \xrightarrow{k_1} Y$$

the value of k_1 is 1×10^{-4} s⁻¹. If the reaction starts from **X**, the ratio of the concentrations of **Y** and **Z** at any given time during the course of the reaction is found to be $[\underline{Y}] = \underline{1}$

The value of k2 is

(A) $1 \times 10^{-4} \mathrm{s}^{-1}$	(B) $2.5 \times 10^{-5} \text{ s}^{-1}$
(C) $4 \times 10^{-4} \text{ s}^{-1}$	(D) $4 \times 10^4 \text{ s}^{-1}$

Q. 21 to Q. 60 carry two marks each.

- Q.21 The correct order of v_{CO} for the compounds [Mo(CO)₃(NMe₃)₃], [Mo(CO)₃(P(OPh)₃)₃], [Mo(CO)₃(PCl₃)₃], [Mo(CO)₃(PCl₃)₃] in the IR spectrum is
 - $(A) [Mo(CO)_3(NMe_3)_3] > [Mo(CO)_3(P(OPh)_3)_3] > [Mo(CO)_3(PMe_3)_3] > [Mo(CO)_3(PCl_3)_3] > [Mo(CO)_3(PCl$
 - $(B) [Mo(CO)_3(PCl_3)_3] > [Mo(CO)_3(NMe_3)_3] > [Mo(CO)_3(P(OPh)_3)_3] > [Mo(CO)_3(PMe_3)_3]$
 - $(C) [Mo(CO)_3(PCl_3)_3] > [Mo(CO)_3(P(OPh)_3)_3] > [Mo(CO)_3(PMe_3)_3] > [Mo(CO)_3(NMe_3)_3]$
 - (D) $[Mo(CO)_3(PMe_3)_3] > [Mo(CO)_3(NMe_3)_3] > [Mo(CO)_3(PCl_3)_3] > [Mo(CO)_3(P(OPh)_3)_3]$
- Q.22 2.5 g of an iron compound upon suitable treatment yielded 0.391 g of iron(III) oxide. The percentage of iron in the compound is

(atomic weight of Fe: 55.847, O: 15.994)

(D) 9.11

(A) 10.94 (B) 12.15 (C) 11.31

Q.23 In the reaction

Ph₃P Mel [X] n-BuLi

the compounds X and Y, respectively, are

- (A) [Ph₃P(Me)I]; Ph₃P=CH–CH₂–CH₂–CH₃
- (C) [Ph₃P(Me)₂]; Ph₃P=CH₂

(B) $[Ph_3P(Me)][I]; Ph_3P=CH_2$

(D) [Ph₃P(Me)][I]; Ph₃P

Q.24 The ¹H NMR spectrum of HD consists of a (A) singlet (B) 1:1 doublet

(C) 1:1:1 triplet (D) 1:2:1

(D) 1:2:1 triplet

6/16

CY

Q.25	The X-ray powder pattern of NaCl shows an intense cone at $\theta = 15.87^{\circ}$ using X-rays of wavelength	ł
	1.54×10^{-8} cm. The spacing between the planes (in Å) of NaCl crystal is	

(A) 1.41 (B) 2.82 (C) 4.23 (D) 5.63

Q.26 Among the following, the isoelectronic and isostructural pair is

(A) CO_2 and SO_2 (B) SO_3 and SeO_3 (C) NO_2^+ and TeO_2 (D) SiO_4^{4-} and PO_4^{3-}

Q.27 Two samples have been given to you : [NiCl₂(PPh₃)₂] and [PdCl₂(PPh₃)₂]. A physical method that can be used to identify these compounds unambiguously is

(A) HPLC	(B) magnetic susceptibility	
(C) ¹³ C NMR spectroscopy	(D) Mössbauer spectroscop	

Q.28 In the reaction $HSO_4^-(aq) + OH^-(aq) \implies SO_4^{2-}(aq) + H_2O(l)$, the conjugate acid-base pairs are

(A) HSO_4^- and SO_4^{2-} ; H_2O and OH^- (C) HSO_4^- and OH^- ; SO_4^{2-} and H_2O (B) HSO_4^- and H_3O^+ ; SO_4^{2-} and OH^- (D) HSO_4^- and OH^- ; SO_4^{2-} and H_3O^+

Q.29 Designate the following complexes **X**, **Y** and **Z** as inert or labile: $\mathbf{X} = [Al(C_2O_4)_3]^{3-}, \quad \mathbf{Y} = [V(H_2O)_6]^{2+}, \quad \mathbf{Z} = [Cr(C_2O_4)_3]^{3-}$

(A) X and Y are inert; Z is labile(C) X is inert; Y and Z are labile

(B) X and Z are labile; Y is inert(D) X is labile; Y and Z are inert

Q.30 In the reaction sequence



X and Y, respectively, are



2009

Q.31 The major product \mathbf{X} (based on the preferred conformation) in the reaction



(Ts = p-toluenesulfonyl)



Q.32 In the reactions

is

 $[X] \xrightarrow{PhCHO} D-glucose \xrightarrow{acetone} [Y]$

the major products X and Y, respectively, are



Q.33 In the reaction



the major product X is

(A)
$$\xrightarrow{D}$$
 (B) \xrightarrow{H} (C) \xrightarrow{D} (C) \xrightarrow{D} (D) \xrightarrow{D} (D) \xrightarrow{D}

8/16

Q.34 Reaction of *m*-methylanisole with lithium in liquid ammonia and *t*-butyl alcohol at -33 °C generates compound X as the major product. Treatment of the compound X with dilute sulphuric acid produces compound Y as the major product. The compounds X and Y, respectively, are

CY

9/16



- Q.35 The number of signals that appear in the broad-band decoupled ¹³C NMR spectrum of *ortho-*, *meta-* and *para-*dichlorobenzenes, respectively, are
 - (A) 3, 4 and 2 (B) 3, 3 and 2 (C) 4, 4 and 2 (D) 3, 4 and 4
- Q.36 In the reaction sequence

 $\frac{Me}{90\%} \xrightarrow{\text{NH}_2\text{OH}} [Y] \xrightarrow{\text{H}_2\text{SO}_4} [Z]$



2009

the structure of the major product Z and the overall yield for its formation from the ketone X, are



Q.37 In the reaction sequence



the major products X and Y, respectively, are



Q.38 In the reaction sequence

$$\frac{PdCl_2, CuCl}{O_2, H_2O}$$
 [X] $\frac{H_2SO_4}{DMF}$ [Y]

the major products X and Y, respectively, are



Q.39 In the reaction sequence

$$Ph' \rightarrow [X] \xrightarrow{\text{conc.}} [Y]$$

the major products X and Y, respectively, are



Q.40 In the photochemical reaction

٨

formation of the compound X can be inferred by the disappearance of the ¹H NMR signal at [¹H NMR spectrum of the starting material : δ 9.7 (1H, s), 7.8 (1H, d, J 8.0 Hz), 7.1 – 6.8 (2H, m), 3.9 (3H, s), 2.5 (3H, s) ppm]

(A)	δ 9.7 ppm	0.0	(B)	δ 7.8 ppm
(C)	δ 3.9 ppm		(D)	δ 2.5 ppm

Q.41 The half-life $(t_{1/2})$ for the hydrolysis of an ester varies with the initial concentration of the reactant ([E]₀) as follows:

$\begin{array}{l} [E]_0 / 10^{-2} \mbox{ mol } L^{-1} \\ t_{1/2} / \mbox{ s} \end{array}$		5.0 240	4.0 300		3.0 400	
The order of the read	tion is					
(A) 0	(B)	1	(C)	2		(D) 3

CY

CY

2009				CY		
Q.42	The fluorescence lifetime of a molecule in solution is 10 ns. If the fluorescence quantum yield is 0.1, the rate constant of fluorescence decay is					
	(A) $1 \times 10^9 \text{s}^{-1}$		(B) $1 \times 10^8 \text{ s}^{-1}$			
	(C) $1 \times 10^7 \text{ s}^{-1}$		(D) $9 \times 10^7 \text{ s}^{-1}$			
	(0) 1110 0		(2) 21120 0			
Q.43	The fundamental vibe The relative population respective ground state	The fundamental vibrational wavenumbers for H_2 and I_2 are 4403.2 cm ⁻¹ and 214.5 cm ⁻¹ , respectively. The relative population of the first excited vibrational states of these two molecules compared to their respective ground states at 300 K are, respectively :				
	(A) 6.75×10^{-1} and 3	57×10^{-1}	(B) 6.75×10^{-10} and	3.57×10^{-1}		
	(C) 3.57×10^{-6} and 6.	75×10^{-1}	(D) 3.57×10^{-1} and 6	5.75×10^{-1}		
	(0) 5.57 A 10 and 0.	2.01	(1) 5.57 1 10 414 0			
Q.44	The degeneracy of a sis	quantum particle in a cu	bic box having energy fo	our times that of the lowest energy		
	(A) 3	(B) 6	(C) 1	(D) 4		
			C.B. In the support phase, phase	Giores the grote friedfold of		
Q.45	The rotational Ramar and 19223.707 cm ⁻¹ .	n spectrum of ¹⁹ F ₂ shows The rotational constant	a series of Stokes lines a for ${}^{19}F_2$ in GHz is	at 19230.769 cm ⁻¹ , 19227.238 cm ⁻¹		
	(A) 26.484	(B) 52.968	(C) 105.936	(D) 3.531		
	()		(0)	Common Data Questions		
Q.46	The de Broglie wave	length for a He atom trav	velling at 1000 ms ⁻¹ (typ	ical speed at room temperature) is		
	(A) 99.7×10^{-12} m	(B) 199.4×10^{-12} m	(C) 199.4×10^{-18} m	(D) 99×10^{-6} m		
0.17	Given that the stan	dard molar anthalaias	of formation of NO/a	and NO (a) are reconciliable		
Q.47	90.3 kJ mol ⁻¹ and 33.2 kJ mol ⁻¹ , the enthalpy change for the reaction $2NO(g) + O_2(g) \rightarrow 2NO_2(g)$ is					
	(A) 16.6 kJ	(B) -57.1 kJ	(C) –114.2 kJ	(D) 57.1 kJ		
			construction and			
Q.48	Among the following	g, the equilibrium which	is NOT affected by an in	ncrease in pressure is		
	$(A) 2SO_3(g) \rightleftharpoons 2SO_3(g)$	$O_2(g) + O_2(g)$				
	(B) $H_2(\alpha) + I_2(\alpha) \rightarrow$	2HI(g)				
	(D) 112(B) + 12(S) -	2111(8)				

- $(C) C(s) + H_2O(g) \rightleftharpoons CO(g) + H_2(g)$
- (D) $3Fe(s) + 4H_2O(g) \Longrightarrow Fe_3O_4(s) + 4H_2(g)$
- Q.49 The free energy change (ΔG) of 1 mole of an ideal gas that is compressed isothermally from 1 atm to 2 atm is

(A) RTln2	(B) -2RT	(C) -RTln2	(D) 2RT
-----------	----------	------------	---------

CY

11/16

Q.50 Two liquids B and C form an ideal solution. In the figure below, the vapour pressure P of this solution is shown as a function of the mole fraction, X_B, of component B.



Given a state of this vapour-liquid mixture whose overall composition corresponds to point E in the figure, the mole fraction of B in the vapour phase is approximately

- (A) 0.25
- (C) 0.65

(B) 0.53 (D) 0.80

Common Data Questions

Common Data for Questions 51 and 52 :

Treatment of $W(CO)_6$ with 1 equivalent of $Na(C_5H_5)$ in THF solution gives the ionic compound M. Reaction of M with glacial acetic acid results in product N. The ¹H NMR spectrum of N displays two singlets of relative intensity 5:1. When N is heated, hydrogen gas is evolved and O is produced; O may also be prepared by refluxing $W(CO)_6$ with cyclopentadiene and H_2 is also produced. Treatment of O with an equivalent of Br_2 produces P. (Use the 18 electron rule as your guide).

Q.51 The compounds M and N, respectively, are

- (A) $[(C_5H_5)W(CO)_3]$ Na and $[(C_5H_5)W(CO)_3H]$
- (B) $[(C_5H_5)W(CO)_4]$ Na and $[(C_5H_5)W(CO)_4H]$
- (C) [(C₅H₅)W(CO)₃]Na and [(C₅H₅)W(CO)₄H]
- (D) $[(C_5H_5)W(CO)_4]$ Na and $[(C_5H_5)W(CO)_3H]$
- Q.52 The compounds O and P, respectively, are

(A) $[(C_5H_5)W(CO)_3]_2$	and	$[(C_5H_5)W(CO)_3Br]$
(B) [(C ₅ H ₅)W(CO) ₄]	and	$[(C_5H_5)W(CO)_2Br(THF)]$
(C) $[(C_5H_5)W(CO)_2(THF)_2]$	and	$[(C_5H_5)W(CO)_3Br]$
(D) $[(C_5H_5)W(CO)_3]_2$	and	[(C ₅ H ₅)W(CO) ₂ Br(THF)]

2009

Common Data for Questions 53 and 54 :

An organic compound X (C₉H₁₀O) exhibited the following spectral data.

 $IR : 1680 \text{ cm}^{-1}$

¹H NMR : δ 7.8 (2 H, d, J 7.5 Hz), 7.2 (2 H, d, J 7.5 Hz), 2.7 (3 H, s) and 2.4 (3 H, s). Compound **X** on treatment with *m*-chloroperbenzoic acid produced two isomeric compounds **Y** (major) and **Z** (minor).

Q.53 Compounds Y and Z, respectively, are



Q.54 Compounds Y and Z can be differentiated by carrying out basic hydrolysis, because

(A) Y produces 4-methylphenol and Z is unaffected

(B) Y produces 4-methylphenol and Z produces 4-methylbenzoic acid

(C) Y is unaffected and Z produces 4-methylbenzoic acid

(D) Y is unaffected and Z produces 4-methylphenol

Common Data for Questions 55 and 56 :

Character table for the point group C2v is given below:

C_{2v}	Е	C ₂	$\sigma_v(xz)$	$\sigma_v(yz)$	- D.	1.
A1	1	1	1	1	Z	x^{2}, y^{2}, z^{2}
A ₂	1	1	-1	-1	Rz	xy
B1	1	-1	1	-1	x, R _v	XZ
B ₂	1	-1	-1	1	y, R _x	yz

Q.55The reducible representation corresponding to the three translational degrees of freedom, Γ_{tr} , is(A) 3, 1, 1, 1(B) 3, -1, 1, 1(C) 3, -1, -1, -1(D) 3, 1, -1, -1

Q.56 The asymmetric stretching mode of the H_2O is shown below. The molecular plane is yz and the symmetry axis of H_2O is z.

This vibration transforms as the irreducible representation

(A) A_1 (B) B_1 (C) A_2 (D) B_2

Statement for Linked Questions 57 and 58 :

Triphosphazene is prepared by reacting X and Y in equimolar ratio at 120-150 °C using appropriate solvents.

- Q.57 The reactants X and Y, respectively, are
 - (A) PCl₃; NH₃
 - (C) PCl5; NH4Cl

(B) PCl₅; NH₃(D) PCl₃; NH₄Cl

Q.58 The structure of triphosphazene is







Statement for Linked Questions 59 and 60 :

In the reaction mechanism given below,

 $X + Y \xrightarrow{k_{1,} E_{A,1}} Z \xrightarrow{k_{3}, E_{A,3}} P$

'k's represent rate constants, 'EA's represent activation energies, and k2 >> k3.

Q.59 The overall rate constant (koverall) for the formation of P can be expressed as

A) k_1k_3 / k_2	(B) k ₁
C) $k_1 / (k_2 + k_3)$	(D) $k_1 / (k_2 - k_3)$

Q.60 The overall activation energy (EA, overall) for the formation of P can be expressed as

(A)
$$\frac{E_{A,1} \cdot E_{A,3}}{E_{A,2}}$$

(B) $E_{A,1}$
(C) $E_{A,1} + E_{A,3} - E_{A,2}$
(D) $\frac{E_{A,1}}{E_{A,2} + E_{A,3}}$