## ALCHEMIST SCIENCE ACADEMY



# Assignments

With Previous Years Questions CSIR-NET Dec 2018, TIFR- 2018 & GATE 2019

## Spectroscopy



for

CSIR NET/JRF, GATE, TIFR, JNU, Ph.D

## ASSIGNMENT

## Spectroscopy

### **ALCHEMIST SCIENCE ACADEMY**

Website: www.csirnetalchemist.com

E-mail: <u>alchemistscienceacademy@gmail.com</u>

Head Office: 28-A Jia Sarai, Hauz Khas New Delhi – 16 Ph. 011-26511021, 8285787633, 9582285416, 9953942156

#### **Contents**

	Page no
MB, ESR, PES, UV	3-27
Mass Spectroscopy	28-33
NMR Spectroscopy	34-75
Molecular Spectroscopy	76-99

### Spectroscopy (M-B, ESR, PES, UV) NET Previous Year's Question

Q1.	Which of the follo	owing spectroscopic tech	niques will be useful to disti	nguish betweer	M - SCN and				
	M - NCS binding	modes?			[NET June 2011]				
	(a) NMR	(b) IR	(c) EPR	(d) Mass					
Q2.	The Stark splitting is true? ( $\mu$ is the $\epsilon$	for a given field is larger dipole moment)	for a molecule AX as compar	ed to BX. Which	one of the following [NET June 2011]				
	(a) $\mu_{\mathrm{AX}} = \mu_{\mathrm{BX}}$	(b) $\mu_{\mathrm{AX}} > \mu_{\mathrm{BX}}$	(c) $\mu_{\rm\scriptscriptstyle AX} < \mu_{\rm\scriptscriptstyle BX}$	(d) $\mu_{_{ m B}}$	x = 24.x				
Q3.	A molecule, AX has	s a vibrational energy of	100 cm <sup>-1</sup> and rotational en	ergy of 10cm	Another molecule				
	BX, has a vibration	BX, has a vibrational energy of 400 cm and rotational energy of 40cm statements about the coupling of vibrational and rotational motion is true							
	(a) The coupling is	stronger in BX							
	(b) The coupling is								
	(c) Magnitude of coupling is same in both AX and BX								
	(d) There is no coup	oling in both AX and BX							
Q4.	The order of carbor	nyl stretching frequency i	in the IR spectra of ketone, a	amide anhydrid	e is:				
					[NET June 2011]				
	(a) Anhydride > ami		(b) Ketone > amide	: > anhydride					
	(c) Amide > anhydri	a sh the	(d) Anhydride > Ket	tone > amide					
Q5.	The absorption at )	$279$ nm ( $\varepsilon = 15$ ) in	the UV spectrum of acetone	is due to	[NET June 2011]				
	(a) $\pi - \pi^*$ transition	on	(b) $n-\pi^*$ transition	on					
	(c) $\sigma - \sigma$ transition	pn e	(d) $\pi - \sigma^*$ transiti	on					
Q6.	In the EPR spectrum	of tetragonal Cu(II) con	nplex, when $\left. \mathrm{g} \right\  \mathrm{gl} > \mathrm{g}_{\mathrm{e}}$ the	e unpaired elec	tron resides in the				
	orbital.		One ce	,	[NET June 2011]				
	(a) q''	(b) $d_{x^2-y^2}$	(c) $d^2$	(d) $d_{\checkmark}$					
Q7.	Consider the Compo	unds,			[NET June 2011]				
	(A) SnF <sub>4</sub> (B) SnCl <sub>4</sub> an	nd (C) R <sub>3</sub> SnO	CI						
	The nuclear quadrupo	ole splitting are observed	d for						
	(a) (A),(B) and (C)	(b) (A) and (B) only	(c) (B) and (C) only	(d) (A) an	d (C) on;y				

The correct value of isomer shift (in Mossbauer spectra) and its exaplanation for Fe(II) – TPP and Fe(III) – Q8. [NET June 2011] TPP respectively from the following are:

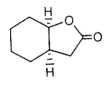
( TPP = tetraphenylporphyrinate)

(A) 0.52mms

- (B) 0.45mms <sup>1</sup>
- (C)Increase in s electron density
- (D) Decrease in s electron density
- (a) (A) and (D); (B) and (C)
- (b) (A) and (C); (B) and (C)
- (c)(B)and (D); (A) and (D)
- (d) (B) and (D); (A) and (C)

In the IR spectrum, carbonyl absorption band for the following compound appears at Q9.

[NET Dec. 2011]



- (a) 1810cm
- (b) 1770cm
- (d) 1690cm

In  $^{57}$  Fe' Mossbauer experiment source of 14.4 keV (equivalent to  $3.48 \times 10^{12} MHz$ ) is moved towards Q10. The shift in frequency of the source for this sample is absorber at a velocity of 2.2mms [NET Dec. 2011]

- (a) 35.5 MHz
- (b)25.5MHz
- (c) 20.2MHz
- (d) 15.5MHz
- If Mossbauer spectrum of Fe(CO), is recorded in the presence of a magnetic field, the original spectrum Q11. [NET June 2012] with two lines changes into the one with
  - (a) Three lines
- (b) Four lines
- (c) Five lines
- (d) Six lines
- The total numbers of fine and hyperfine EPR lines expected for octahedral high spin Mn(II) Complexes are [NET June 2012] respectively (1 = 5/2 for Mn)
  - (a) 3 and 30
- (b) 5 and 33
- (c) 5 and 30
- (d) 4 and 24
- The number of lines exhibited by a high resolution EPR spectrum of the species, O13.

[NET Dec. 2012]

[Cu(ethylenediamine)2]2 + is [Nuclearspin (I) of Cu = 3/2 and that of N = 1]

- (a) 12
- (b) 15
- (c) 20
- (d)36

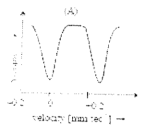
Q14. In the IR spectrum of p-nitrophenyl acetate, the carbonyl absorption band appears at [NET Dec. 2012]

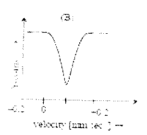
- (a) 1670 cm <sup>--1</sup>
- (b) 1700 cm -1
- (c) 1730 cm
- (d) 1760 cm

Q15. The recoil energy of a Massabauer nuclide of mass 139 amu is 2.5MeV. The energy emitted by the nucleus in KeV is [NET Dec. 2012]

- (a) 12.5
- (b) 15.0
- (c) 2.05
- (d) 25.0

Q16. The Mossbuar spectra of two iron complexes are shown below. They may arise from (i) high-spin iron(III), (ii) high-spin iron(III) and (iii) low-spin iron(III) The correct matches of spectra (A) and (B) with the iron complexes are





- (a) A with (i) and B with (ii)
- (b) A with (ii) and B with (i)
- (c) A with (iii) and B with (ii)
- (d) A with (ii) and B with (iii)

Q17. The number of EPR signals observed for octahedral Ni(II) complexes is

[NET June 2013]

- (a) One
- (b) Two
- (c) Three
- (d) Zero

Q18. A triatomic molecule of the type AB, shows two IR absorption lines and one IR-Raman line. The structure of the molecule is [NET June 2013]

(a) B - B - A

(b) B - A - B





Q19 In the IR, spectrum, the absorption band due to carbonyl group in phenyl acetate appears at

[NET June 2013]

- (a) 1800 cm <sup>-1</sup>
- (b) 1760 cm<sup>-1</sup>
- (c) 1710 cm
- (d) 1660 cm <sup>1</sup>

Q20. Among the following, those can act as Mossbauer nuclei are

[NET June 2013]

- (A)  $^{129}$ l
- (B) 57 CO
- (C) 57 Fe
- (D) 121 Sb

(a) A, B, C and D

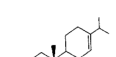
(b) B,C and D only

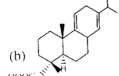
(c) A,B and D only

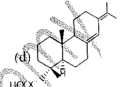
(d) A,C and D only

- For a tetragonally distorted Cr(III) complex, zero field splitting results in the following number of Kramers Q21. doublets:
  - (a) 1

- (b) 2
- (c) 3
- (d) 4
- In the UV-V is spectrum, a diterpenoid exhibited a  $\lambda_{\rm max}$  at 275 nm. The compound, among the choice given Q22. [NET June 2013] below is







In the IR spectrum of p-nitrophenyl acetate, the carbonyl absorption band appears at Q23.

[NET Dec. 2013]

- (a) 1660 cm
- (b) 1700 cm<sup>-1</sup>
- (c) 1730 cm
- (d) 1770 cm <sup>-1</sup>
- In the atomic absorption spectroscopic estimation of Fe(III) using  $O_2/H_2$  flame, the absorbance decreases Q24. [NET Dec. 2013] with the addition of
  - (a)  $CO_3^{2n}$
- (c) EDTA
- The number of lines in the ESR spectrum of  $\mbox{CD}_3$  is (the spin of D is l ) Q25.

[NET June 2014]

(a) I

- (c) 4
- (d)7
- The correct order of the isomeric shift in Mossbauer spectra ( <sup>57</sup> Fe source) of iron compound is Q26.

[NET Dec. 2014]

(b) Fe(III) > Fe(II) > FE(IV)

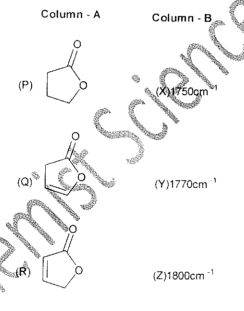
Fe (III) > Fe(II)

- (d) Fe(IV) > FE(II) > FE (III)
- $m{A}_{c}$  compound with molecular formula  $C_{_4} H_{_6} O_{_2}$  shows band at 1770 cm $^{-1}$  in IR spectrum and peaks at 178,68,28 and 22 ppm in  $^{13}$  CNMR  $^{13}$  spectrum. The correct structure of the compound is Q27. [NET Dec. 2014]





Q28. The gas commonly used in generating plasma in inductively Coupled Plasma Atomic Emission spectroscopy (ICPAES) is [NET Dec. 2014] (a) Argon (b) Carbon dioxide (c) Nitrous oxide (d) hydrogen Q29. The compound that exhibits sharp bands at 3300 and 2150 cm<sup>-1</sup> in the IR spectrum is [NET June 2015] (a) 1 - butyne (b) 2 - butyne (c) butyronitrile (d) butylamine The reduced form of a metal ion M in a complex is NMR active. On oxidation, the complex gives an EPR Q30. signal with  $\,{
m g} \approx 2.2$  and  $\,g_{\perp} \approx 2.0$  . Mossbauer spectroscopy cannot characteristic the metal complex. The M is [NET June 2015] (a) Zn (b) Sn (c) Cu Q31. The resonance Raman stretching frequency  $(v_{n-n}, \text{ in cm}^{-1})$  of  $O_2$  is 1580 The voxyhemoglobin in close to [NET Dec. 2015] (a) 1600 (c) 800 (d) 1100 Q32. Correctly matched structure and carbonyl stretching frequency set, i [NET Dec. 2015]



(a) P - Y, Q - Z, R - X

(b) P - Y, Q - X, R - Z

(c) P - Z, Q - Y, R - X

(d) P - X, Q - Z, R - Y

Q33. Pick the correct statements about Atomic Absorption Spectrometry (AAS) from the following

[NET Dec. 2015]

- (A) Hg lamp is not a suitable source for AAS
- (B) Graphite furnace is the best atomizer for AAS
- (C) Non-metals cannot be determined with AAS
- (D) AAS is better than ICP-AES for simultaneous determination of metal ions. Correct answer is

(a) A,B and C

(b) B,C and D

(c) A,B and D

(d) B and D

Q34. Mossbauer spectrum of a metal complex gives information about

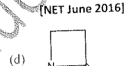
[NET Dec. 2015]

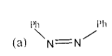
- (A) oxidation state and spin state of metal
- (B) types of lingands coordinated to metal
- (C) nuclear spin state of metal
- (d) geometry of metal

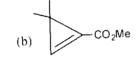
Correct answer is

- (a) A and C
- (b) B and C
- (c) A, B and D
- (d) B and D

Q35. Among the following, the compound that displays an IR band at 2150 cm<sup>-1</sup> is









- Q36. In the UV-visible absorption spectrum of an  $\alpha-\beta$  unsaturated carbonyl compound, with increasing solvent polarity,
  - (a)  $n-\pi^*$  transitions undergo hypsochromic shift,  $\pi-\pi^*$  undergo bathochromic shift
  - (b)  $n-\pi^*$  transitions undergo bathochromic shift,  $\pi-\pi^*$  undergo hypsochromic shift
  - (c) both  $n-\pi^*$  and  $\pi-\pi_*$  transition undergo bathochromic shift
  - (d) both  $n-\pi^*$  and  $\pi^*$  transitions undergo hypsochromic shift.
- Q37. Identify correct statements for the EPR spectrum of VO  $\left(acac\right)_2$  [with square pyramidal geometry at vanadium] at  $77K\left[\left(\sqrt[4]{V}\right)=7/2\right]$ .
  - (A) It has two g values

(B) It has 8 lines only

(C) It has one g value

(D) It has two patterns of  $8\,$  lines each.

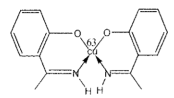
- (a) A and D
- (b) A and C
- (c) B and C
- (d) B and D
- Q38. The record Mossbauer spectrum of Fe containing samples, a source, 'X' is used. X after a nuclear transformation (Y) gives  $\lambda$  radiation used in Mossbauer spectroscopy. [NET June 2016]
  - (a)  $^{57}$  Fe,  $\beta$  emission
- (b)  $^{57}$ Co, $\beta$  wmission

(c) 57 Co, e capture

(d) 57 Fe,e capture

- Q39. The compound which shows IR frequencies both 3314 and 2126  $cm^{\pm 1}$  is
- [NET Dec. 2016]

- (a)  $CH_3 (CH_2)_4 CH_2 SH$
- (b)  $CH_3(CH_2)_4 CH_2C = N$
- (c)  $CH_3 (CH_2)_4 CH_2 C = C H$
- (d)  $CH_1(CH_2)_1 C = C(CH_2)_2 CH_3$
- Q40. For complex A, deuteration of NH protons does not alter the EPR spectrum. The number of hyperfine line expected in the EPR  $\left|I\left({}^{63}Cu\right)=\frac{3}{2}\right|$  spectrum of A is [NET Dec. 2016]



- (a) 20
- (b) 12
- (c) 60
- (d) 36
- Q41. Mossbauer spectrum of complex [Fe(I, I0 phenanthroline)] (NCS), show two line at 300 K, four lines at 186K and again two lines at 77K. This can be attributed to [NET June 2017]
  - (A) Change in the coordination mode of NCS
  - (B) Change in the spin-state of iron
  - (C) cis-trans isomerisation
  - (D) change in metal-ligand bond distances

The correct statements are

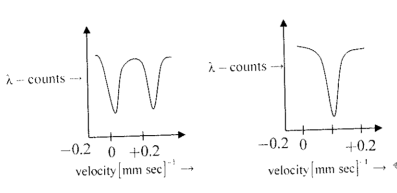
- (a) A and B
- (b) B and C
- (c) A and C
- (d) B and D
- Q42.  $(R_3Ge)_2$  on photolysis gives a radical which shows ESR spectrum. The ESR signals carrying the signature of  $^{73}Ge(1=9/2)$  are in terms of [NET June 2017]
  - (a) Nine line
- (b) Tenlines
- (c) two lines
- (d) one line
- Q43. The resonance Raman stretching frequency  $(in \ cm^{-1})$  of the bound  $O_2$  species in oxy-hemerthyrin and oxyhemoglobin, respectively, are [NET June 2017]
  - (a) ~850 and 1100

(b) ~ 750 and 850

(c) ~ 850 and 850

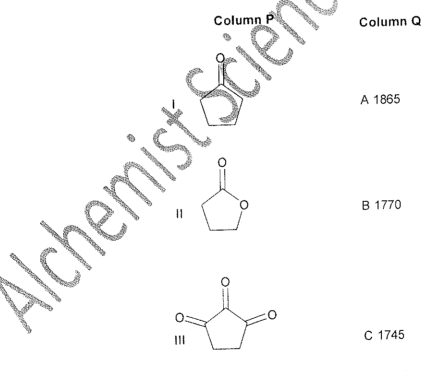
(d) ~ 1100 and 850

Q44. The Mossbaur spectra of two iron complexes are shown below. The may arise from (i) high spin iron(III) (ii) high-spin iron (II) and (iii) low-spin iron(III)



The correct matches of spectra (A) and (B) with the iron complex are

- (a) A with (i) and B with (ii)
- (b) A with (ii) and B with (i)
- (c) A with (iii) and B with (ii)
- (d) A with (ii) and B with (iii)
- Q45. Correct match of the compounds in column P with the IR stretching frequencies (cm<sup>-1</sup>) in Column Q is [Net Dec. 2018]



- (a) I B; II C; III A
- (b) I C; II A; III B
- (c) I C; II B; III A
- (d) 1 A; II C; III B

Q46. The number of lines in EPR spectrum of  ${\rm CD_3}({\rm I_{D+1}})$ 

[NET June 2018]

(a) 3

- (b) 5
- (c) 7

(d) 9

- Consider the species  $No, I_2, I_2^+, Cu^{2+}$  and  $VO^{2-}$ . The number of paramagnetic species among them and Q47. the EPR inactive species, respectively, are
  - (a) 4 an 1,
- (b) 4 and I,
- (c) 3 and  $VO^{2+}$ ,  $Cu^{2+}$  (d) 3 and NO,  $Cu^{2+}$
- In IR spectra the stretching frequency (in  $\,\mathrm{cm}^{-1}$ ) of the carbonyl group of the following compounds is in Q48. the order [NET Dec. 2018]







- (a) B > A > C
- (b) A > C > B
- (c) B > C > A
- Arrange the following molecules in order of increasing fundamental frequencies Q49.

[NET Dec. 2018]

- (a)  $O_2^2 < O_2 < O_2 < O_2^2$
- (c)  $O_2^2 < O_2 < O_2^+ < O_2^-$

#### Answer Key

Q1.(b)	2.(b)	3.(a)	4.(d)	5.(c)	6.(b)	7.(d)
8.(a)	9.(b)	10.(b)	11.(d)	12.(c)	13.(d)	14.(d)
15.(d)	16.(a)	10.(b) 17.(b)	18.(b)	19.(d)	20.(b)	21.(c)
22.(d) C	23 (b)	24.(d)	25.(a)	26.(a)	27.(c)	28.(a)
29.(a)	30.(c)	31.(d)	32.(a)	33.(a)	34.(d)	35.(c)
36.(a)	37.(c)	38.(c)	39.(c)	40.(a)	41.(d)	42.(b)
43.(d) A	44.(b)	45.(c)	46.(c)	47.(b)	48.(c)	49(a)

#### **GATE Previous Year's Question**

Q1. Among the isomers of  $C_4H_6$  given below, the compound which exhibits an babsorption band at 3300cm

in the IR spectrum, is

[GATE 2002]

- (a) 1,3 butadiene
- (b) 1 butyne
- (c) 2 butyne
- (d) cyclobutene.

Q2.

On the basis of Woodward-Fieser rules, the dienes that have  $k_{
m max}$  values in the range 268 – 273nm are

[GATE 2003]

- (a) P and Q
- (b) P and R 🌸
- (c) Q and R
- (d) Q and S
- Q3. The number of hyperfine split lines observed in ESR spectrum of methyl radical is

[GATE 2004]

(a) 1

(h) 4

(c) 6

- (d) 8
- Q4. Match the compounds P  $\sim$  S with their carbonyl stretching frequency  $(cm^{-1})I VI$  in IR spectroscopy.

[GATE 2004]

P. acetone

- I. 1870
- Q. ethyl acetate
- II. 1800

- R. acetamide
- III. 1740
- **s**. acetyl chloride
- IV.1700
- V.1660
- VI. 1600
- (a) P IV, Q III, R I, S VI
- (b) P-III, Q-VI, R-V, S-II
- (c) P = IV, Q = III, R = V, S = II
- (d) P-II, Q-V, R-III, S-VI

(a) 1-(i), 2-(ii), 3-(iii), 4-(iv)

(c) 1-(ii), 2-(i), 3-(iv), 4-(iii)

Q5. Q6. Q7.	(a) tungaten filament (c) deuterium lamp The moleculer active  (a) CO <sub>2</sub> A radical contains <sup>14</sup> hyperfine constant 0.  (a) 3 line  Which of the following	in rotational micr (b) $SF_6$ N(I=1) with hy 35mT. The ESR sp (b) 6 line	not an excitation source for (b) Nernst glower (d) mercury are rowave, infrared absorption (c) HCI perfine constant 1.61mT an ectrum will exhibit. (c) 7 line	as well as rotational $\mbox{(d)} \ \ \mbox{H}_2$	[GATE 2006]
	The moleculer active  (a) CO <sub>2</sub> A radical contains <sup>14</sup> the hyperfine constant 0.  (a) 3 line  Which of the following	(b) $\operatorname{SF}_6$ N $(1=1)$ with hy 35mT. The ESR sp	owave, infrared absorption (c) HCI perfine constant 1.61mT an ectrum will exhibit.	(d) H <sub>2</sub> nd Two equivalent pr	[GATE 2006] otons (I = 1/2) with
	(a) CO <sub>2</sub> A radical contains <sup>14</sup> th hyperfine constant 0.  (a) 3 line  Which of the following	(b) $\operatorname{SF}_6$ N $(1=1)$ with hy 35mT. The ESR sp	(c) HCl perfine constant 1.61mT an ectrum will exhibit.	(d) H <sub>2</sub> nd Two equivalent pr	[GATE 2006] otons (I = 1/2) with
Q7.	A radical contains 14 phyperfine constant 0.  (a) 3 line  Which of the following	$N\left(I=I\right)$ with hy 35mT. The ESR sp	perfine constant 1.61mT an ectrum will exhibit.	ad Two equivalent pr	otons (I = 1/2) witl
Q7.	hyperfine constant 0.  (a) 3 line  Which of the following	35mT. The ESR sp (b) 6 line	ectrum will exhibit.		1
	Which of the followin		(c) 7 line	(d) 9 line	
		a absorptions is s			
Q8.		e molar absorptiv			ectrum recorded in
	(a) $\lambda_{\max} 217 \text{nm} \left( \epsilon_{\max} =$		(b) $\lambda_{\text{max}} 214 \text{nm} (\epsilon_{\text{max}})$		
	(c) $\lambda_{\text{max}} 253 \text{nm} \left( \epsilon_{\text{max}} =$	= 50,000)	(d) $\lambda_{max} 250 \text{nm} \left(\epsilon_{max}\right)$	= 500)	
Q9.	Match of the compou	nds in <b>list-I</b> with t	the stretching frequencies	on of the princip	ole function groups
	given in List-II				[GATE 2006]
		(1) (2) (3)	(ii) 2224 (iii) 179 (iii) 179	95	
		(4)	N (iv) 17	'50	
			(v) 16	95	
	(a) 1-ill, 2-iv, 3-l, 4-v		(b) 1-iii, 2-iv, 3-ii, 4-v		
4	(c) 1-iv, 2-v, 3-ii, 4-i		(d) 1-iv, 2-iii, 3-v, 4-i		
Q10.	Match the observed pr absorption in list-II	inciple absorption	ns in the visible spectrum sh	own in <b>List-</b> ł With th	e bond shows this [GATE 2007]
	List-I	List-II			
	$(1) \ \sigma \to \sigma^*$	(i) C – C			
	$(2) n \to \sigma^*$	(ii) C – O			
	(3) $n, \pi^*$	(iii) C = O			
	$(4) - \pi , -\pi^*$	(iv) C = C			

(b) 1-(i), 2-(iii), 3-(ii), 4-(iv)

(d) 1-(iv), 2-(ii), 3-(iii), 4-(i)

Q11. The IR stretching frequencies (cm ') for the compound X are as follows:3300 – 3500 (s,br);3000(m);

2225(s);1680(s)

[GATE 2008]

The correct assignment of the absorption band is

(a) 
$$\overline{v}_{(\text{CH})} = 3300 - 3500; \overline{v}_{(\text{CH})} = 3000; \overline{v}_{(\text{CN})} = 2225; \overline{v}_{(\text{CO})} = 1680$$

(b) 
$$\overline{v}_{(\text{OH})} = 3300; \overline{v}_{(\text{CH})} = 3000 - 3500; \overline{v}_{(\text{CN})} = 2225; \overline{v}_{(\text{CO})} = 1680$$

(c) 
$$\overline{v}_{(\text{OH})} = 3300 - 3500; \overline{v}_{(\text{CH})} = 3000; \overline{v}_{(\text{CN})} = 1680; \overline{v}_{(\text{CO})} = 2225$$

(a) 
$$\overline{v}_{(\text{OH})} = 3300; \overline{v}_{(\text{CH})} = 3300 - 3500; \overline{v}_{(\text{CN})} = 1680; \overline{v}_{(\text{CO})} = 2225$$

- Q12. The total number of ways in which two nonidentical spin 1/2 particles can be oriented relative to a constant magnetic fields is; [GATE 2008]
  - (a) 1

- (b) 2
- (c) 3
- (d) 4
- Q13. The Oxidation sate of the metal ion in the catalyst can be deducted by
- [GATE 2008]

- (a) Atomic absorption spectroscopy
- (b)Mossbauer spectroscopy

(c) HPLC

(d) Gas Chromatography

#### Common data for Q 45 and Q. 16

A six –coordinate transition-metal complex is ESR and Mossbauer active. The effective magnetic moment of this complex is 5.9 B.M.

- Q14. The metal ion along with its oxidation state and the number of unpaired electron present are [GATE 2010]
  - (a) Fe(II) and 4

(b) Mn(II) and 5

(c) Fe(Ni) and 1

(d) Fe(III) and 5

Q15. The complex is

[GATE 2010]

(a)  $\left[ Mn \left( H_2O \right)_6 \right]^2$ 

(b)  $\left[ \text{Fe} \left( \text{CN} \right)_{6} \right]^{3}$ 

(c)  $\left[ \text{Fe} \left( \text{H}_2 \text{O} \right)_6 \right]^{2-}$ 

(d)  $\left[ \text{Fe} \left( \text{H}_2 \text{O} \right)_6 \right]^{3+}$ 

Q16. The extent of Mossbauer quadrupole splitting of iron follows the order

[GATE 2012]

(a) 
$$\operatorname{FeCl}_2 4H_2O > k_2 \left[ \operatorname{Fe}(CN)_s (NO) \right] > \operatorname{FeCl}_3 6H_2O$$

(b) 
$$K_2[Fe(CN)_s(NO)] > FeCl_2.4H_2O > FeCl_3.6H_2O$$

(c) 
$$FeCl_1.6H_2O > K_2[Fe(CN)_2(NO)] > FeCl_2.4H_2O$$

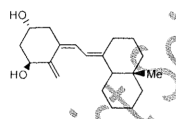
(d) 
$$FeCl_2.4H_2O > FeCl_3.6H_2O > K_2[Fe(CN)_s(NO)]$$

- Q17. Among the compounds given in the option (a) to (d), the one that exhibits a sharp band at around 3300 cm<sup>-1</sup> In the IR spectrum is [GATE 2014]
  - (a) 1,2 butadiene

(b) 1, 3 - butadiene

(c) 1 - butyne

- (d) 2- butyne
- Q18. Given the fact 1,3 butadiene has a UV absorption of 217 nm, the absorption wavelength (in nm) for the conjugated system shown below is [GATE 2014]



(Use these absorption values for auxochromic groups:

Alkyl: +5 exo-cyclic double bond +5 every additional conjugated C = C 30)

Q19. In atomic absorption spectroscopy, the atomization process utilizes

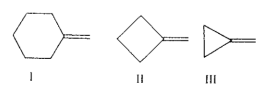
[GATE 2014]

(a) flame

(b) electric field

(c) magnetic field

- (d) electron beam
- Q20. The correct order of IR stretching frequency of the C = C in the following olefins is [GATE 2014]



- (a) I > II > III
- (b) || > ||| > |
- (c) ||| > || > |
- (d)||| > || > ||

- Q21. The value of 'g' and the number of signal observed for the reference standard diphenylpicrylhydrazyl (DPPH), in the solid state ESR spectrum are, respectively [GATE 2015]
  - (a) 2.0036 and 1

(b) 2.0036 and 3

(c) 2.2416 and 1

- (d) 2.2416 and 3
- Q22. The  $v_{0-0}$  resonance resonance Raman stretching frequency (in cm $^{-1}$ ) of the  $O_2$  coordinated to iron centre in oxyhemoglobin is nearly
  - (a) 1100
- (b) 850
- (c) 1550
- (d) 1950
- Q23. The characteristic feature of an electron spin resonance (ESR) Spectrum of frozen agueous solution of CuSO<sub>4</sub>.5H<sub>2</sub>O at 77 k is [GATE 2017]
  - (a)  $g_{\parallel} > g_{\perp}$

22.(a)

- (b)  $g_{\parallel} < g_{\perp}$
- (c)  $g_{\parallel} = g$
- (d)  $g_x \neq g_y \neq g_z$

#### Answer Key

7.(d) 6.(c) 5.(d) 2.(d) Q1.(b) 14.(d) 13.(b) 12.(a) 11.(a) 9.(c) 8.(a) 21.(a) 20.(c) 19.(a) 18.(282) 16.(d) 15.(d)

> Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

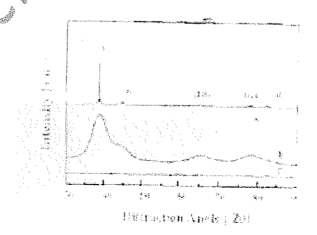
#### **TIFR Previous Year's Question**

- Q1. It is not easy to locate H atoms in the structure of proteins obtained by X-ray diffraction due to which of the following reasons: [TIFR 2010]
  - (a) H atoms are transparent to the X-rays
  - (b) H bonding interaction present in protein structures
  - (c) interaction of H with solvent molecules
  - (d) Very low levels of scattering of X-rays by H
- Q2. Arrange the characteristic timescalses of the following set of dynamical spectroscopic techniques in decreasing order of time (longest to shortest); NMR (Nuclear Magnetic Resonance), ESR (Electron spin Resonance), Fluorescence, Raman and absorption.
  - (a) absorption > NMR > Fluorescence > ESR > Raman
  - (b) NMR > ESR > Fluorescence > Raman > Absorption
  - (c) Fluorescence > Absorption > ESR > NMR > Raman
  - (d) All have similar characteristic timescales.
- Q3. In a chemistry lab, the aim of an experiment was to generate well structured, 5 nm sized gold nanoparticles.

  A student experimented with the following synthesis: [TIFR 2013]

I-nonanethiol  $(C_9H_{19}SH)$  and HAuCl were first mixed (in a molar ratio of 10:1, and then gold ions were reduced by slowly adding NaBH in this synthesis after 2 hours no precipitate was observed and only a very faint pink colloidal solution was obtained. This solution was then evaporated onto a glass slide and the sample characterized by X-ray diffraction.

Given below are three XRD patterns. One but of these three patterns was obtained by the student when he / she characterized the sample on the glass slides. Which statement below is true?



Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

- (a) The synthesis did not yield gold nanoparticles as there was no precipitate, therefore the XRD pattern that will be obtained is that shown as curve (c) It is almost a flat baseline indicating no product.
- (b) 5 nm gold nanoparticles were formed and XRD pattern shown as (a) represents the product. It signifies that the nanoparticles are crystalline and the face-centered cubic (fcc) crystal structure can be clearly used to index the peaks.
- (c) 5nm gold nanoparticles will show size-dependent line broadening and therefore if the product consisted of such particles, then curve (b) will be obtained.
- (d) None of the above statements are true.
- Q4. What are the limits of detection of the following common analytical methods used with capillary separations:

Fluorescence, mass spectrometry, UV-Vis absorbance, and NMR, respectively, in mole [TIFE 2013]

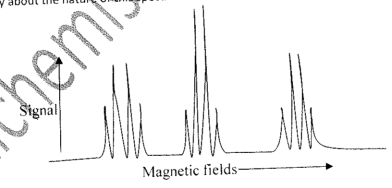
(a) 
$$10^{18} - 10^{-23}$$
,  $10^{-13} - 10^{-21}$ ,  $10^{13} - 10^{-16}$ ,  $10^{-9} - 10^{-1}$ 

(b) 
$$10^{-13} - 10^{-21}$$
,  $10^{-18} - 10^{-23}$ ,  $10^{-13} - 10^{-16}$ ,  $10^{-9}$ 

(c) 
$$10^{-18} - 10^{-23} \cdot 10^{-13} - 10^{-21} \cdot 10^{-9} = 10^{-11} \cdot 10^{-16} - 10^{-16}$$

(d) 
$$10^{-13} - 10^{-21}, 10^{-13} - 10^{-13}, 10^{-13}, 10^{-9}, 10^{-9}, 10^{-9}$$

Q5. A magnetic resonance spectrum, recorded using a radiation of frequency 100 is shown below. What can you say about the nature of this spectrum?



- (a) It is a continuous-wave NMR spectrum
- (b) It is fourier transform NMR spectrum
- (c) It is an ESR spectrum
- (d) None of the above

Q6. A long column of water in any transparent bottle appears slightly blue. However, If we replace water with

heavy water  $\left(D_2O\right)$  it will look more transparent. This effect is due to

[TIFR 2016]

- (a) Rayleigh scattering
- (b) Kinetic isotope effects
- (c) Absorption spectra of H<sub>2</sub>O and D<sub>2</sub>O are different
- (d) None of the above.
- Q7. For electronics transitions in organic molecules, the expected energy ordering the transitions is:

TIFR 2018)

- (a)  $\pi$  to  $\pi^*$  in to  $\sigma^* < \sigma$  to  $\sigma^* < n$  to  $\pi^*$
- (b)  $\pi$  to  $\pi^* < n$  to  $\pi^* < n$  to  $\sigma^* < \sigma$  to  $\sigma^*$
- (c) n to  $\pi^*$   $\pi$  to  $\pi^* <$  n to  $\sigma^* <$   $\sigma$  to  $\sigma^*$
- (d) n to  $\sigma^* < \sigma^*$  to  $\sigma^* < n$  to  $\pi^* < \pi$  to  $\pi^*$

#### Answer Key

Q1.(\*)

2.(\*)

3.(c)

4 (3)

<sup>₩</sup> 5.(c

6.(c)

7.(c)

### Other Examinations Previous Year's Question

Q1.	The two characteristic stretch	ning frequencies $(cm^{-1})$ observed in	the IR spectrum of compounds
Q1.	containing NO <sub>2</sub> group is		
	(a) 3400 and 3300	(b) 1860 and 1760	
	(c) 1550 and 1350	(d) 2250 and 1760	
Q2.	The UV-light source used in UV	-visible spectrophotometer is	
	(a) Mercury lamp	(b) Tungsten lamp	
	(c) Deuterium lamp	(d) Sodium lamp	
Q3.	The recoil velocity ( in ms <sup>-1</sup> ) o	f a free Mossbauer nucleus of mass 10 2	kg while emitting a γ - ray of 0.1
	nm wavelength is:		
	(a) 6.626 (b) 66.26	(c) 662.6 (D) 6626	
Q4.	The number of lines that appe	ar in the EPR spectrum of $\left[ \mathbf{C_6} \mathbf{\hat{H}}_6  ight]$ is	
	(a) 5 (b)		(d) 13
Q5.	Which of the following molect	ules will not absorb infrared radiation?	
	(a) CO (b)	(c) COCl <sub>2</sub>	(d) CO <sub>2</sub>
Q6.	Which of the following molec	ules will have the highest zero point vibra	itional energy?
-		CH <sub>4</sub> (c) CCl <sub>4</sub>	(d) CBr <sub>4</sub>
Q7.	For CO molecule.		
4	(a) All the vibrational modes	are either IR or Raman active	
		nan active vibrations will be the same	
	(c) All vibrations are IR active		
	(d) All vibrations are Raman a	active.	the following
Q8.	The presence of hydrogen bo technique	anding in an organic compound can easily	pe established dulizing the rolls and
	(a) IR spectroscopy	(b) Mass spectra	
	(c) Cyclic voltametry	(d) CD-ORD	
		- 11 1	44001¢

Q9. Which of the following will most conveniently confirm if a know solid sample is impure?

- (a) NMR
- (b) Mass spectrum
- (c) IR spectrum
- (d) Melting point

Q10. The molecule azulene has an absorption maximum at 700nm, the red and of the visible spectrum. The next shortest wavelength occurs at 357 nm. The predicated color that azulene can exhibit is:

- (a) blue
- (b) red
- (c) indigo
- (d) green

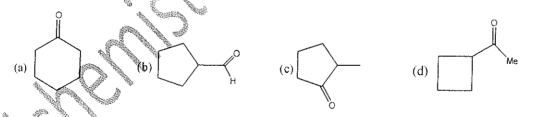
Q11. The anti Stokes lines are generally weaker in intensity because, the

- (a) molecules absorb the radiation completely
- (b) incident radiation is unpolarized
- (c) atoms are generally in the ground state:
- (d) vibrational energies are small.

Q12. Which of the two compounds will be most readily separated by TLC (thin layer chromatography)?

- (a) naphthalene and anthracene
- (b) naphthalene and acetophenone
- (c) acetophenone and 4- methylacetophenone
- (d) benzoic acid and 3- toluic acid.

Q13. Compound X having molecular formula CHOO exhibits absorption at 1700cm in the IR spectrum and forms oxime. Compound X yields adipic acid on oxidation. The structure of the compound X is:



Q14 What is the  $\lambda_{\rm max}$  for the following compound? Use the provided parameters for your calculation.

Transoid base value 214nm

Cisoid base value 253nm

Alkyl groups on base + 5nm

Extanded condiation + 30 nm

- (a) 234nm
- (b) 244nm
- (c) 273nm
- (d) 283nm

- Of the molecules  $\mathrm{CH_4},\mathrm{CO_2}$  , benzene and  $\mathrm{H_2}$  , the ones that will absorb infrared radiation are Q15.
  - (a) CH<sub>4</sub>, CO<sub>2</sub> benzene
- (b) CH<sub>a</sub>, benzene and H,
- (c)  $CO_2$  benzene and  $H_2$
- (d)  $\mathrm{CH_4}, \mathrm{CO_2}\,\mathrm{and}\ \mathrm{H_2}$
- Which one of the normal modes of ethylene is active in the infared? Q16.









- The H-H distance in  $H_{\rm S}$  molecule can be determine by, Q17.
  - (a) Microwave rotational spectroscopy
- (b) Infrared vibrational spectroscopy

(b) NMR spectroscopy

- (d) Rotational Raman spectroscopy
- The virbrational stretching frequency of  $N_2$  can be determined using Q18.
  - (a) Infrared spectroscopy
- (b) Microwaye spectroscopy
- (c) Raman spectroscopy
- (d) NMR spectroscopy
- The number of IR active vibrational modes in ammonia is: Q19.
  - (a) 6

(c) 2

- (d)3
- will not have anyabsorbance in the microwave or the infrared region of the The molecule that Q20. electromagnetic spectrum is:
- (b) HF
- (c) CH<sub>4</sub>
- (d)  $H_2$
- the infrared spectrum of  $\mathrm{CO}_2$  exhibits the following number of absorptions: Q21.
  - (a) One
- (b) Two
- (c) Three
- (d) Four

- Which of the following molecules shows EPR resonance? Q22.
  - (a) H<sub>2</sub>O
- (b) O.
- (c) H,O,
- (d) CO<sub>2</sub>

- Q23. Which of the following electronics transitions is disallowed?
  - (a)  $\pi \rightarrow \pi^*$
- (b)  $\sigma \rightarrow \sigma^*$
- (c)  $n \longrightarrow \pi^*$
- (d)  $\delta \rightarrow \delta^*$
- Q24. The order of  $\lambda_{max}$  in the UV-Visible spectra for compound. A C is:



Δ



В



(a) A > B > C

(b) B > A > C

(c) B > C > A

(d) C > B > A

- Q25. The number of vibrational degrees of freedom in a gaseous mixture of 30 CI 0.37 CI 0 and 30 CI 0.17 CI 0 is
  - (a) 1
- (b) 2
- (c) 3

- (d) 4
- Q26. IR stretching frequency of carbonyl group in aldehydes are acid chlorides in  $cm^{-1}$  are
  - (a) 1730 1700 and 1650 1580
- (b) 1680 1660 and 1730 1700
- (c) 1730 1700 and 1820 1770
- (d) 1680 1660 and 1820 1770
- Q27. The number of peaks in the ESR spectrum of CH<sub>3</sub> radical is:
  - (a) 1

- b) 2
- (c)3

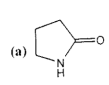
- (d) 4
- Q28. In the UV spectrum of cyclohexenone, the absorption at  $\lambda_{\rm max}$  ~215nm is due to the transition
  - (a)  $\sigma \rightarrow \sigma'$
- (b)  $\sigma \rightarrow \iota$
- (c)  $\pi \rightarrow n$
- (d)  $\pi \rightarrow \pi^*$
- Q29. The most convenient spectroscopic technique to establish the presence of inter-molecular hydrogen bonding in hydroxy compounds is
  - (a) UV
- (b) IR
- (c) EPR
- (d) Mass

- Q30. Spectrum of a gaseous containing  $H_1D_2O_2$  and  $N_2$  shows
  - (a) 4150, 1560, 2200 and 2950
- (b) 1560, 2200, 4150 and 2950
- (c) 4150, 2950, 1560 and 2200
- (d) 2950, 2200, 1560 and 4150

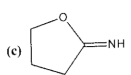
Alche	Tillst Science From Transfer F							
Q31.	The source of ultra-violet radiation	used in UV-visible Spectrophotometer is						
	(a) Mercury vapour lamp	(b) Sodium vapour lamp						
	(c) Halogen vapour lamp	(d) Hydrogen vapour lamp.						
Q32.	Among the following, the correct s							
	(a) Absorption of radiation shifts to	longer wavelength region with increase in conjugation						
	(b) Absorption of radiation shifts to higher energy region with increase in conjugation							
	(c) Intensity of $n \to \pi^*$ transition	decreases upon conjugation.						
	(d) Intensity of $\sigma  ightarrow \sigma^{st}$ transition	n decreases upon conjugation.						
Q33.	The infrared and Raman spectrum	of BF <sub>3</sub> are expected to show						
	(a) The same number of peaks							
	(b) More absorption peaks in IR in comparison to Raman							
	(c) More absorption peaks in Ram							
	(d) Absorption peaks present in R							
Q34.	Among the following diatomic mo	lecutes, the one that shows EPR signals is:						
	(a) Li <sub>2</sub> (b) B <sub>3</sub>	(c) C <sub>2</sub> (d) N <sub>2</sub>						
Q35.	Which of the following molecule	, will have $n o\pi^*$ transition at the longest wavelength ?						
	(a) HCHO	(b) $CH_3COC_2H_5$						
504	(c) C.H.COC.H.	(d) CH <sub>3</sub> COC <sub>6</sub> H <sub>5</sub>						
Q36.	The total number of vibrational o	egrees of freedom of $H_2 {\cal O}_2$ is:						
	(a) 7 (b) 6	(c) 4 (d) 9						
Q37.	Resonant frequencies for EPR an	d NMR are respectively in the spectral region						
	(a) Microwave and Far-IR	(b) Far –IR and microwave						
	(c) Radiofrequency and microwa	ve (d) Microwave and radiofrequency						

Q38.	Two samples have	e been given to you:	$\left[\operatorname{NiCl}_{2}\left(\operatorname{PPh}_{2}\right)_{2}\right]$ and $\left[\operatorname{PdCl}\right]$	$\left( \text{PPh}_{3} \right)_{2}$ . A physical method that c	an				
		be used to identify these compounds unambiguously is							
	(a) HPLC		(b) Magnetic suscepti	bility					
	(c) <sup>13</sup> C NMR spe	ctroscopy	(d) Mossbauer spectro	оѕсору					
Q39.	Which among the	following electronic tr	ansitions will have the low						
				<u>-</u> .					
	(a) $n \rightarrow \sigma^*$	(b) $n \rightarrow \pi^*$	(c) $\sigma \longrightarrow \sigma^*$	(d) $\pi \to \pi^*$					
Q40.	Which of the follow	wing molecules shows	s rotational absorption spe	ectrum?					
	(a) CO <sub>2</sub>	(b) OCS	(c) CH <sub>4</sub>	(d) C <sub>2</sub> H,					
Q41.	The H-H distance in	$_{1}~\mathrm{H_{2}}$ molecule can de	etermined by,						
	(a) Microwave rota	tional spectroscopy	(b) Infrared vi	br <b>atio</b> nal spectroscopy					
	(c) NMR spectrosco	рру	(d) Rotational	Raman spectroscopy					
Q42.	The number of vibr	ational degrees of fre	edom in a gaseous mixture	of <sup>35</sup> Cl <sub>2</sub> O, <sup>37</sup> Cl <sub>2</sub> O and <sup>35</sup> Cl <sup>37</sup> ClO is					
	(a) 1	(b) 2	( <b>c</b> ) 3	(d) 4					
Q43.	Cis and Transe cinn	amic acids can be mos	st readily <b>distinguished</b> and	didentified by,					
	(a) IR spectra	à	(b) UV spectra						
	(c) Chemical shift of	f the olefinic hydroger	(d) Coupling co	onstant of the olefinic hydrogens.					
Q44.	The region of electr	omagnetic spectrum	employed in the electron s	pin resonance (ESR) spectroscopy is					
	(a) radiowave	(b) microwave	(c) infrared	/ althoraches					
	(a) radiowave	(b) incrowave	(c) iiii aieu	(d) visible					
Q45.	The number of rota	tional degrees of freed	dom of $\mathrm{CO}_2$ is						
	(a) one	(b) two	(c) three	(d) four					
Q46.	Identify the molecul	e whose rotational co	nstant can not be determi	ned by spectroscopic methods					
4	(a) GFT	(b) H <sub>2</sub>	(c) CO <sub>2</sub>	(d) HCL					
Q4 <b>7</b> .	Obs <b>erve th</b> e following	ng statements							
			d from rotational spectra						
			rotational spectral lines o	f gaseons NO is 2B cm					
				from TMS using a NMR spectrometer					
		. Its chemical shift is 3		,					
	Which of the following	ng is correct?							
	(a) I,II and Illare corre	ect	(b) only III is correct						
	(c) I and II are correct	:	(d) Only I is correct						

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416 Q48. Which among the following exhibits a carbonyl absorbtion band at 1770cm









- Q49. Among the following molecules, one having the highest zero point vibrational energy is
  - (a) HF
- (b) CH<sub>1</sub>
- (c) H,O
- (d) NH<sub>3</sub>
- Q50. The number of hyperfine lines in the EPR spectrum of a one electron reduced product of  $\left[\text{Co}_3\left(\text{CO}\right)_9\,\text{Se}\right]$  (I = 7/2 for Co nucleus) is
  - (a) 8

- (b) 15
- (c) 22
- (d) 1

- Q51. Which of the followings species is ESR active?
  - (a) VOSO<sub>4</sub>
- (b) K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>
- (c) KMnO<sub>4</sub>
- (d)  $\left[\operatorname{Co}\left(\operatorname{NH}_{3}\right)_{6}\right]\operatorname{Cl}$

- Q52. Which one of the following exhibits rotational spectra?
  - (a) H<sub>2</sub>
- (b) N
- 🏿 (c) CO
- (d) CO<sub>2</sub>
- Q53. The bond that gives the most intense band in the infrared spectrum for its stretching vibrations is
  - (a) C H
- (b) N H
- (c) 0 H
- (d) S H
- Q54. An examination of saturated hydrocarbons containing methyl group show asymmetrical  $(V_{,})$  Stretching modes in the region of
  - (a) 2960 and 2870 cm
- (b) 3200 and 3100  $\,cm^{-1}$

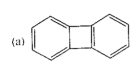
(c) 1800 - 1700 cm

(d) 1650 - 1450 cm<sup>-1</sup>

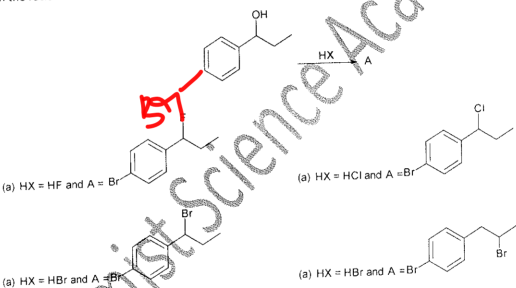
			Answer Ke	У		
1.(c)	2.(c)	3.(b)	4.(b)	5.(b)	6.(b)	7.(a)
8.(a)	9.(d)	10.(d)	11.(c)	12.(b)	13.(a)	14.(d)
15.(a)	16.(c)	17.(d)	18.(c)	19.(a)	20.(d)	21.(c)
22.(b)	23.(c)	24.(b)	25.(c)	26.(c)	27.(b)	28.(d)
29.(b)	30.(c)	31.(a)	32.(a)	33.(c)	34.(b)	35. <b>(c</b> )
36.(b)	37.(d)	38.(d)	39.(b)	40.(b)	41 (d)	42.(c)
43.(b)	44.(b)	45.(b)	46.(b)	47.(c)	48.(a)	49.(b)
50.(c)	51.(d)	52.(c)	53.(c)	<b>54.</b> (a)		

### Mass Spectroscopy NET Previous Year's Question

- Q1. In the mass spectrum of dodecahedrane  $(C_{20}H_{20})$ , approximate ratio of the peaks at m / z 260 and 261 [NET Dec. 2011]
  - (a) 1:1
- (b) 5:1
- (c) 10:1
- (d) 20:1
- Q2. Anthranilic acid, on treatment with iso-amyl nitrite furnishes a product which displays a strong peak at 76(m/e) in its mass spectrum. The structure. The structure of the product is [NET June 2014]



Q3. The mass spectrum of the product A, formed in the following reaction, exhibits M, M + 2, M + 4 peaks in the ratio of about 1 : 2 : 1. The reagent HX and the product A are [NET June 2014]



Q4. In The mass of metastable ion produced due to decomposition of  $F_l^+$  in the following mass fragmentation sequence is

Diethyal-phthalate  $\longrightarrow$   $F_1^+$   $\longrightarrow$   $F_2^+$  + CO (d) 210.2

Q5. In the mass spectrum of the compound given below, during the  $\alpha-$  cleavage, the order of preferential loss of groups is [NET June 2011]

Me OH 
$$C_3H_7$$

- (a) Me  $> C_3H_7 > Et$
- (b)  $C_3H_7 > \text{Et} > \text{Me}$
- (c) Et > Me > C<sub>3</sub>H<sub>2</sub>
- (d) Et  $> C_3H_7 > Me$

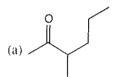
- Q6. In the mass spectrum of 1,2 dichloroethane, approximate ratio of peaks at m/z value 98,100,102 will be [NET Dec. 2015]
  - (a) 3:1:1
- (b) 9:6:1
- (c) 1:1:2
- (d) 1: 2:1
- Q7. Mass fragment of  $[IrCI]^{+}$  in mass spectrometry shows three mass peaks at m/z = 226,228 and 230. Given that natural abundances of  $^{-191}$  Ir.  $^{-193}$  Ir.  $^{-35}$  Cl and  $^{-37}$  Cl are 37%, 63%, 76% and 24% respectively, the intensities of the mass peaks in the order [Net June 2017]
  - (a) 49.5:100:26.6

(b) 100:49.5:26.6

(c) 26.6:100:49.5

- (d) 26.6:49.5:100
- Q8. Among the following, the compound that gives base peak at m/z 72 in the El mass spectrum is

[NET Dec. 2017]



- (b)
- (c)
- (d)
- Q9. Mass spectrum of a compound shows an [M+2] for peak that is about 4% of  $M^{\circ}$ . This indicates that the compound has one.
  - (a) Fluorine
- (b) sulfur
- (c) bromine
- (d) chlorine

#### **Answer Key**

- 1. (b)
- 2.(b)
- 3. (c)
- 4. (b)
- 5. (b)
- 6.(b)
- 7.(a)

3.(a)

#### **GATE Previous Year's Question**

Statement: Bromopyrimidine  $(C_4H_3BrN_2)$  exhibits two prominent peaks in the mass spectrum at m/z 158 Q1. and 160 in the ratio 1:1

Reason: There are two basic centres in the molecule, which are protonated.

Assertion: There are two isotopes of bromine,  $^{79}$  Br and  $^{81}$  Br , that occur in the ratio of 1 : 1

[GATE 2004]

Choose the correct answer from the following four choices.

- (a) Both Reason and assertion are correct.
- (b)Both Reason and Assertion are wrong
- (c) Reason is correct and Assertion is wrong
- (d) Reason is wrong but Assertion is correct
- Among the isomers  $C_{i0}H_{i4}$  shown Q2.

[GATE 2007]

The isomer that can be identified uniquely by mass spectrometry alone is

- (a) P
- (c) R

(d) S

Linked Answer Type Q3. And Q.4

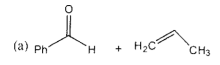
For butyrophenone ( $PhCOCH_3CH_3CH_3$ )

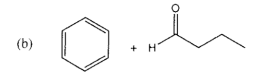
The most probable fragmentation observed in the electron impact ionization (EI) mass spectrometry is Q3.

(d) Ph 
$$H + H_2C$$
  $CH_3$ 

Q4. Photirradiation leads to the following set of products:

[GATE 2008]

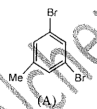




- Q5. The ratio of relative intensities of the two molecular ion peaks of methyl bromide (CH, Br) in the mass spectrum is [GATE 2012]
  - (a)  $M^+: (M+2) = 1.3$
- (b)  $M^+:(M+2)^+=3$
- (c)  $M^+: (M+2)^+ = 1:1$
- (d)  $M^+:(M+2)^+=1:2$
- Q6. The m/z value of the detectable fragment formed by Lafferty like rearrangement of the following compound in mass spectrometer is [GATE: 2014]



Q7. The mass spectrum of a dihalo compound shows peaks with relative intensities of 1 : 2 : 1 Corresponding to M, M + 2 and M + 4 (M is the mass of the molecular ion), respectively. The compound is [GATE 2015]



(B)





(D)

(6)

(a) A

(b) B

(c) C

(C)

- (d) D
- Q8. In the electron ionization (EI) mass spectra, methyl hexanoate, methyl heptanoate and methyl octanoate give.

  The same base peak. The m/z value of the base peak is [GATE2018]

#### **Answer Key**

1. (d)

- 2.(b)
- 3. (a)
- 4. (d)
- 5. (c)
- 6.(41)
- 7.(a)

8.(72)

#### **TIFR Previous Year's Question**

1. (c)	2.(b)	3. (a)	4. (đ)	5. (*)	6.(*) <b>C</b>
			Answ	ver Key	
	(a) 1 – bromopropan	e (b) 2 – bron	nopropane	(c) Both A and B	(d)) None of these
	124. Large fragment i	ions are seen at m	n/z = 43 (base	peak)	[TIFR 2019]
Q6.	Identify the compour	nd whose molecul	lar ion appear	s as a pair of equal ir	intensity peaks at $m/z = 122 \& m/z$
	(a) 2.24 and 0.0126	(b) 50 and 2		(c) 2.24 and 0.025	(d) 1.12 and 0.0126
					[TIFR 2019]
	the peaks at mass nu	mber 139and 140	. Isotopic abui	ndances: <sup>13</sup> C, 98.89 <sup>6</sup>	% <sup>13</sup> C,1.11%, <sup>19</sup> F,100%
Q5.	If the peak in the ma	ss spectrum of C	$_2\mathrm{F}_6^{}$ at mass nu	ımber 138 is 100 uni	it tall, what will be the heights of
	(c) The highest mass r	earran <b>ge</b> mention		(d) The ion peak gre	
	(a) The molecular ion	45		(b) The lowest m/z p	
Q4.	Which of the following	g statements is th			in a mass spectrum? [TIFR 2016]
	(c) 50 and 25			(d) 2.24 and 0.025	
	(a) 2.24 and 0.0126			(b) 1.12 and 0.0126	
	( Isotopic abundances	: ¹² C, 98.89% ; ¹³ )	C.1.11%; ''F	A STATE OF THE STA	
	the peaks at mass nun	ibei 133 and 140	•		
Q3.	If the peak in the mass	s spectrum of $C_2$	${\sf F}_{\!\scriptscriptstyle 6}$ at mass nu	mber 138 is 100 unit	s tall, what will be the heights of [TIFR 2015]
	(a) 1252	(b) 1374		(c) 1498	(d) 1325
	The approximate m/z	value of the conse			vould be
		2016, 1767, 1			
Q2.	Electrospray ionization this technique to hem the protein. The three	eprotein it gave a consecutive peak	a large numbe s are observed	er of beaks corspond	is of proteins. On application of ing to different charge-states of [TIFR 2013]
	(a) $^{79}\mathrm{Br}$ and $^{80}\mathrm{Br}$				
	numbers 158,160 and	(b) $^{80}$ Br and		c) $^{70}\mathrm{Br}$ and $^{81}\mathrm{Br}$	(d) $^{79}\mathrm{Br.}^{80}\mathrm{Br}$ and $^{81}\mathrm{Br}$
	numbers 158 160 and	162. Which isotor	es of bromine	Occur in nature :	[TIFR 2010]

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

#### Other Examination Previous Year's Question

Q1.	The ratio of M	and M + 1 peak	s in the mass spectr	um of ${\sf C}_{60}$ is		
	(a) 1 : 1	(b) 60:1	(c) 3 : 2	(d) 1:60		
Q2.			opes, Br - 79 and Br spectrum of bromot	· 81 that occur in an a penzene?	pproximate 1 :1 ra	atio. What would
	(a) A large pea	k at a m/z value	of 78.			
	(b) A large pea	ık at a m/z value	of 77 and only one r	nolecular ion peak.		
	(c) A large pea	k at m/z value of	77 and two molecu	ar ion peaks of differ	ng heights	
	(d) A large pea	k at a m/z value	of 77 and two moled	cular ion peaks of the	same <b>he</b> ig <b>ht</b>	
Q3.	The base peak	in the electron in	npact mass spectrui	m (El MS) of acetophe	none is	
	(a) 120	(b) 105	(c) 77		(d) 65	
Q4.			oform, the ratio of p nd 81 in equal abud	beaks at $m/2.250$ 2 ance, will be	52 254 ,256 , as	suming bromide
	(a) 1:1:1:1	(b) 1	: 2 : 2 :1	(c) 1 3 3 : 1	(d) 1:6:6	:1
Q5.	15 In the mass	spectrum of dich	lorobenzene the ra	ijo of the peaks at m /	z 146, 148 and 1	50, is
	(a) 1:1:1	(b) 3		(c) 1: 2:1	(d) 9 : 6 : 1	
Q6.	Which one of to (a) $C_2H_3CH(CH)$			Il give a fragment ion	at m/z = 58 in the	ir mass spectra?
	(c) CH <sub>3</sub> CH <sub>2</sub> CH		E.	H <sub>3</sub> ) <sub>2</sub> CHCH <sub>2</sub> CHO		
Q7.	Benzene and be	e <b>nzene</b> - d <sup>a</sup> (hexa	* ideuterated benzen:	e) may be readily dist	inguished by:	
	(a) Thin layer ch			visible (absorption) sp		
	(c) <sup>13</sup> C N M R s	spectroscopy	(d)Mass	spectrometry		
Q8.	Which techniqu	ે ie would you use	to quickly distingui	sh methyl benzoate fi	rom phenyl acetai	te?
	Which technique (a) HNMR (c) Vapor pressu		(b)Mass	spectrometry		
	(c) Vapor pressu	ure osmometry	(d) Elen	nental analysis.		
		\$\tag{\text{\text{\$\sigma}\tex	Answ	er Key		
1. (c)	2.(d)	3. (b)	4. (c)	5. (d)	6.(a)	7.(d)
3. (b)						

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

#### **NMR Spectroscopy NET Previous Year's Question**

An organic compound  $\left(C_{\gamma}H_{\gamma_{2}}O_{\gamma}\right)$  exhibited the following data in the  $^{+}H$  NMR spectrum. Q1.

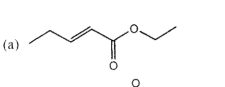
[NET June 2011]

$$\delta 7.10(1H, dt, J = 16 \text{ and } 7.2Hz), 5.90(1H, dt, J = 16 \text{ and } 2Hz)$$

$$4.1(2H,q,J=7.2Hz), 2.10(2H,m), 1.25(3H,t,J=7.2Hz)$$

0.90(3H, t, J = 7.2Hz)ppm

The compound, among the choices given below, is









In the broad decoupled 13 C NMR spectrum, the number of signals appearing for the bicyclooctane A-C Q2. [NET June 2011] respectively, are



(C)

(a) Five, four and eight

(b) Three, two and Five

(c) Five, four and five

- (d) Three, two and eight
- NMR spectrum of PF, the number of singles and multiplicity, at room temperature are Q3.

[NET Dec 2011]

One, singlet

(b) One, doublet

c) two, doublet

- (d) two singlet
- An organic compound  $\left(MF:C_{\mathbf{x}}H_{\mathbf{m}}O\right)$  exhibited the following TH NMR spectral data: Q4.  $\delta 2.5 (3H,s), 3.8 (3H,s), 6.8 (2H,d,J8Hz), 7.2 (2H,d,J8Hz) \text{ ppm. The compound among the choices, is}$ [NET Dec. 2011]
  - (a) 4 ethylphenol

(b) 2 - ethylphenol

(c) 4 -methylanisole

(d) 4- methylbenzył alcohol

Q5. The NMR spectrum of AX<sub>3</sub> exhibits lines at  $\delta = 2.1$  and 2.3ppm (for A type protons), and  $\delta = 4.1, 4.3, 4.5$  and 4.7 ppm (for A type protons), measured from TMS with an instrument operating at 100MHz. The chemical shift (in ppm) of A and X protons and cuping constant (in Hz) are respectively.

[NET Dec. 2011]

(a) 4.4,2.2 and 20

(b) 2,2,4.4 and 10

(c) 2.2,4.4 and 5

(d) 4.3,2.1 and 20

Q6. An organic compound having molecular formula  $C_{15}H_{14}O$  exhibited the following  ${}^{\dagger}H$  and  ${}^{13}C$  NMR spectral data.

<sup>1</sup> H NMR: $\delta 2.4(s)$ , 7.2(d, J = 8Hz), 7.7(d, J = 8Hz)

 $^{13}$ C NMR:  $\delta$ 21.0,129,0.130.0,136.0,141.0,190.0

(b)

- Q7. Appropriate  $^{\dagger}H$  NMR chemical shifts  $\delta$  ) for the protons A-D for the following compound are

[NET Dec. 2011]

(a) A - 6.8, B - 5.7; C - 3.9;D - 2.1ppm

(b) A - 6.8; B - 5.7; C - 2.1; D - 3.9 ppm

(c) A = 5.7, B - 6.8; C - 3.9; D - 2.1 ppm

(d) A - 5.7; B - 6.8; C - 1; D - 3.9 ppm

Q8. A compound A having the composition  $FeC_0H_RO_3$  shown one signal at 2.5 ppm and another one around 5.0 ppm in its  $^3H$  NMR spectrum. The IR spectrum of this compound shows two bands around the 1680 cm  $^{-1}$ . The compound follows the 18 electron rule of the following statements for A, the correct one is/are [NET Dec. 2011]

(A) It has  $\eta^s$  — Cp group

(B) It has terminal CO ligand

(C) It has a CH, ligand

(D) It has Fe - H bond

(a) (A) and (B) only

(b) (A) and (C) only

(c) (A) and (C) only

(d) (B) and (D) only

- The uncertainty in the NMR frequency of a compound in liquid state (relaxation time = 1s) 0.1 Hz Q9. uncertainty in the frequency (in Hz) of same compound in solid state (relaxation time = [NET Dec.2011]
  - (a)  $10^{-4}$
- (b) 100
- (c) 1000
- (d)  $10^{-3}$
- The number of distinct peak in the proton decoupled  $^{13}$  C NMR spectra of the following compounds I III, Q10. [NET June 2012] respectively, are:



(A)

- - **(B)**
- - (C)

- (a) 4.6.8
- (b) 3.4.5
- (d) 2,4.6
- HNMR spectra, The expected chemical shifts are Compounds A and B exhibit two singlets, each in their Q11.

at  $\delta$ 

[NET June 2012]

- 7.7 and 3.9 for B (a) 6.9 and 2 1 for A;
- (b) 7.7 and 3.9 for A; 6.9 and 2.1 for B

(B)

- (c) 6.9 and 3.9 for A; 7.7 and 2.1 for B
- (d) 7.7 and 2.1 for A; 6.9 and 3.9 for B

Match of the following Q12.

[NET June 2012]

Compound

 $^{13}$ C NMR chemical shift ( $\delta ppm$  )

(i) 95

(Ă) Acetic acid

(ii) 115

(B) Acetonitrile (C) Acetone

- (iii) 175
- (D) Carbon tetrachloride
- (iv) 205
- (a) (A) (iii), (B) (ii), (C) (iv), (D) (i)
- (b) (A) (iii), (B) (iv), (C) (i), (D) (ii)
- (c) (A) (i),(B) (ii), (C) (iv), (D) (ii)
- (d) (A) (iii), (B) (i), (C) (iii), (D) (iv)

Q13. In the 400 MHz  $^{\prime}$  HNMR spectrum of organic compound exhibited a doublet. The two lines of the doublet are at  $\delta 2.35$  and 2.38~ppm. The coupling constant (J) value is [NET June 2012]

- (a) 3Hz
- (b) 6Hz
- (c) 9Hz
- (d) 12Hz

Q14. The number of single that appear in the appear in the broadband decoupled <sup>13</sup>C NMR spectrum of phenanthrene and anthracene, respectively are [NET June 2012]

(a) ten and four

(b) ten and ten

(c) sevan and four

(d) sevan and seven

Q15. The carbonyl resonance in C NMR spectrum of

 $\left[\left(\eta^{5}-C_{5}H_{5}\right)Rh\left(CO\right)\right]_{3}\left({}^{108}Rh,nucleaspin,I=1/2,100\%}\right)$  shows a triplet at 65°C owing to the presence of [NET June 2012]

- (a) Terminal CO
- (b) μ<sub>s</sub> -- CO
- (c)  $\mu_s$  CC
- (d)  $\eta^{i} = CH$ .

Q16. The metal complex that exhibits a triplet as well doublet in its P NMR spectrum is [NET June 2012]

- (a) mer  $-[IrCl, (PPh_3)_2]$
- (b) trans = [IrCl(CO)(PPh,), |

(c) fac- $\left[IrCl_3\left(PPh_3\right)_3\right]$ 

(d) [Ir(**P**Ph<sub>1</sub>)]

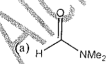
Q17. 'H NMR spectrum of HD would shows

[NET Dec.2012]

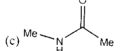
(a) a singlet

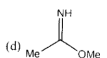
- (b) a doublet
- (c) a triplet with intensity ratio 1:2:1
- (d) a triplet with intensity ratio 1:1:1

Q18. In the 'H NMR spectrum recorded at 293K, an organic compound  $(C_3H_3NO)$ , exhibited singnals at  $\delta 7.8(1H_3S) = 8(3H_3S)$  and  $2.6(3H_3S)$ . The compound is [NET Dec. 2012]



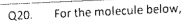
(b) Me NMe2

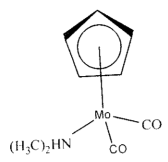




Q19. Consider the following statements for [18] annulene

- (A) It is aromatic
- (B) The inner protons resonate at  $\,\delta$  9.28 in its  $^{1}H$  NMR  $^{2}$  spectrum
- (C) There are six protons in the shielded zone.
- (a) A,B,C
- (b) A and only
- (c) B and C only
- (d) A and C only





Consider the following statements about its room temperature spectral data  $\mathbf{k}$ 

- (A) 1H NMR has singlets at 5.48 and 3.18 ppm
- (B) 1H NMR has multiplet at 5.48 and singlet at 3.18 ppm
- (C) IR has CO stretching bands at 1950 and 1860 cm  $^{\circ}$
- (D) IR has only CO stretching band at 1900 cm<sup>-1</sup>

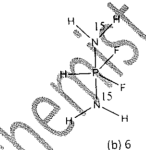
The correct pair of statements is

- (a) A and C
- (b) B and C
- c) A and D

(d) B and D

P NMR singhal for Find out the number lines in the Q21.





(c) 18

(d) 90

- Hydroxybenzoic acid exhibited signals at  $\,\delta$  171,162,133,122  $\,$  and 116 ppm in its broadband decoupoled [NET Dec. 2012]
  - NMR spectrum. The correct assignment of the signals is
  - (a)  $\delta$  171(C 4), 162 (COOH), 133( C 3&5), 122 (C 1) and 116(C 2 & 6) (b)  $\delta$  171 (COOH), 162 (C - 4), 133 ( C - 2 & 6) 122 (C - 1)and 116 (C - 3 & 5)
  - (c) & 171 (C 4), 162 (COOH), 133 (C 2 & 6) (C 1) and 116 (C 3 & 5)
  - (d)  $\delta$  171 (COO), 162 (C 4), 133 (C 3 & 5), 122 (C 1) and 116(C 2 & 6

Q23. An organic compound  $(C_0H_{10}O_3)$  exhibited the following spectral data

[NET Dec.2012]

IR: 34000,180cm<sup>-1</sup>

 $^{1}$ H NMR :  $\delta$ 7.8(IH,d,J = 8Hz),7.0(IH,g,J = 8Hz),6.5(IH,s),5.8(IH,s,D<sub>2</sub>O) exchange geable,

3.9(3H,s), 2.3(3H,s)

The compound is

$$(a) \qquad (b) \qquad (b) \qquad (c) \qquad (d) \qquad (d)$$

- Q24. An AX system gave 4 lines at 4.72, 4.6, 1.12 and 1.0 ppm away from the **JMS** using an NMR spectrometer operating at 100MHz. What are the values of  $J_{AX}$  ( in Hz ) and  $\delta_{AX}$  ( in ppm) respectively [NET June 2013]
  - (a) 12 and 3.6
- (b) 6 and 3.6
- (c) 12 and 2,86
- (d) 6 and 2.86

Q25. Among the following, the correct statements for the following reaction is

[NET June 2013]

$$\frac{1.\text{MeMgBr.EtzQ}}{2.\text{H}_3\text{PO}_4} + \frac{1.\text{MeMgBr.EtzQ}}{(A)}$$

- (a) A is major product and it will have five signals the proton decoupled <sup>13</sup>C NMR spectrum
- (b) A is the minor product and it will have eight signals in the proton decoupled <sup>13</sup>C NMR spectrum
- (c)B is the major product and it will have five signals in the proton decoupled <sup>13</sup>C NMR spectrum
- (d) B is the minocoroduct and it will have five signals in the proton decopled <sup>13</sup>C NMR spectrum
- Q26. In the broad band decoupled <sup>13</sup>C NMR spectrum the number of signals appearing for the two pyrendiols [NET June 2013]

(a) eight and eight

(b) eight and sixteen

(c) five and ten

(d) Five and eight

Q27. An organic compound exhibited the following <sup>1</sup>H NMR spectra data [NET June 2013] & 7.80 (2H,d,J = 8 Hz), 6.80 (2H, d, J = 8Hz), 4.10(2H, q, J = 7.2 Hz) 2.4(3H,s),1.25 (3H, t, J = 7.2Hz)

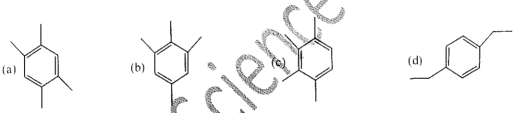
The compound, among the choice given below is

$$(a) \qquad (b) \qquad (c) \qquad (d) \qquad (d) \qquad (e) \qquad (d) \qquad (e) \qquad (e)$$

- Q28. In NMR spectroscopy the product of the nuclear 'g' factor (g), the nuclear magneton ( $\beta$ ) and the magnetic field strength ( $B_0$ ) gives the
  - (a) energy of transition from  $\alpha$  to  $\beta$  state
- (b) Chemical shift

(c) spin-spin coupling constant

- (d) megnetogyric\_ratio
- Q29. A organic compound having the molecular formula  $C_{10}H_{14}$  exhibited two singlets in the  $^{1}H$  NMR spectrum. And three signals in the  $^{13}C$  NMR spectrum. The compound is [NET Dec. 2013]



Q30. Reaction of Ph<sub>2</sub>PCH<sub>2</sub>CH<sub>2</sub>PPh<sub>2</sub> with  $\left[\text{RhCf(CO)}_2\right]_2$  in a 2 : 1 molar ratio gives a crystalline solid A. The IR spectrum of complex A shows  $V_{60}$  at 1985 cm<sup>-1</sup>. The  $^{31}P(^1H)$  NMR spectrum of A consists of two doublets of doublets of equal intensities  $V_{60}$  Rh is 100% abundant and  $V_{60}$ . The structure of complex A is [NET Dec. 2013]

(a) 
$$P$$
 CO  $P$  Ph<sub>2</sub>  $P$ 

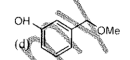
(c) 
$$Ph_2$$
  $CI$   $Ph_2$   $Ph_2$ 

Q31. An organic compound  $(C_6H_{\rm to}O_2)$  , which does not change the color of ferric chloride Solution, exhibited the following H NMR spectral data: [NET Dec. 2013]

 $\delta$  7.3 (1H, t, J = 8Hz), 7.0(1H, d, J = 8 Hz), 6.95(1H,s), 6.9(1H,d,J = 8Hz)

5.3 (1H, brs,  $D_2O$  exachangeable), 4.6 (H, s),3.9 (3H,s). Structure of the compound is





Methyl -4 oxopentanoate exchibited signals at  $\,\delta\,$  208, 172, 51, 37, 32 and 27 ppm in its Q32. CANMR spectrum. The signalsdue to the methoxy, C1, C4 and C5 carbons are [NET Dec. 2016]

(a) OMe - 32; Cl - 208; C4 - 172; C5 - 51

(b) OMe - 51; CI - 208; C4 - 172; C5 - 32

(c) OMe - 32; CI - 172; C4 - 208; C5 - 51

(d) OMe - 51; Cf - 172, C4 - 208; C5 - 32

Given  $\gamma(^{1}H) \approx 2.7 \times 10^{8} \, T^{-1} s^{-1}$  the resonance frequency of a proton in magnetic field of 126 T is close to Q33.

 $(\pi = 3.14)$ 

[NET Dec.2013]

(a) 60 MHz

(b) 110 MHz

🦸 (c) 540 MHz

(d) 780 MHz

The correct match of the  $^1$ H NMR chemical shift  $(\delta)$  of the following species compounds is Q34.

[NET June 2014]







(III)

5.4; IL: 7.2; III : 9.2

(b) I: 9: 2; II: 7.2; III: 5.4

(c) 1 9.2; II : 5.4; III : 7.2

(d) 1: 7.2; II: 9.2, II: 5.4

Q35. The <sup>19</sup>F NMR spectrum of CIF<sub>3</sub> shows

[NET June 2014]

- (a) doublet and triplet for a T-shaped structure
- (b) singlet s for a trigonal planar structure
- (c) singlet for a trigonal pyramidal structure
- (d) doublet and singlet for a T-shaped structure

Q36. The correct  $^{13}C$  NMR chemical  $(\delta)$  shift values of carbons labeled a-e in the following ester are

[NET June 2014]

- (a) a: 19; b: 143; C: 167; d: 125; e: 52
- (b) a:52; b:143; c:167; d:125; e:19
- (c) a : 52; b : 167; c : 143; d : 125 : e : 19
- (d) a: 52; b: 167; c: 125; d: 143; e: 19
- Q37. An organic compound gives following spectral data

[NET June 2014]

IR: 2210,1724cm  $^{-1}$ . H NMR: 1.4(t, J = 7.1Hz, 3H), 4.4(q, J = 7.1Hz, 2H); C NMR: 516,62118,119,125,127,168 the compound is

Q38. The ratio of the relative intensities of the carbon signals in the first order <sup>13</sup> CNMR spectrum of CD<sub>3</sub>Cl is

[NET Dec. 2014]

(a)1:4:6:4:1

- (b) 1:3:3:1
- (c) 1:6:15:20:15:6:1
- (d) 1:3:6:7:6:3:1
- Q39. The following reactions gives a product (recemic) which exhibits the following NMR data [NET Dec. 2014]  $^{\dagger}$ H NMR: $\delta 2.67(3H,s).5.60(2H,s)$

ppm, 13 C NMR: (170.3, 129.0, 105.0, 25.4 ppm

The structure of the product (racemic) is

Q40. An organic compound having molecular formula  $C_{10}H_{12}O_2$  exhibits the following spectral data [NET Dec. 2014]

IR: 3400(br),1600cm<sup>-1</sup>

<sup>1</sup>H NMR:  $\delta 1.85(3H, d, J = 6Hz)$ , 3.8(3H, s),  $5.0(1H, s, D_2O)$  exchangeable)

6.0(1H, dq, J = 18, 6Hz)

6.28 (1H, d, J = 18 Hz), 6.75 (1H, d, J = 8 Hz), 6.8 (1H, s), 6.90 (1H, d, J = 8 Hz)ppm

 $^{13}$ C NMR:  $\delta$ 146.5,144.0,131.0,130.5,123.0,119.0,114.0,108.0,55.0,18.0 ppm

The structure of the compound is

(a) 
$$_{\text{HO}}$$
 (b)  $_{\text{OMe}}$  (b)  $_{\text{OMe}}$  (c)  $_{\text{OMe}}$  (d)  $_{\text{OH}}$  (d)  $_{\text{OH}}$ 

- Q41. A borane (X) is reacted with ammonia to give a salt of borohydride (Y). The <sup>11</sup>B NMR spectrum of Y consists of a triplet and quintet. The borane X is [NET Dec. 2014]
  - (a) B,H,
- (b) B,H,
- (c) B.H
- (d) B<sub>s</sub>H<sub>a</sub>
- Q42. <sup>1</sup>H NMR spectrum of free benzene shows a peak at ~7.2 ppm. The expected chemical shift ( in ppm). The expected chemical shift ( in ppm ) of  $C_nH_n$  ligand in  $^1H$  NMR spectrum of  $\left[\left(\eta^n-C_nH_n\right)Cr(CO)\right]$  and the reason for it, if an, is/are [NET Dec. 2014]
  - (a) 4.5; disruption of ring current
  - (b) 9.0 inductive effect
  - (c) 7.2
  - (d) 2.5 combination of inductive effect and disruption of ring current
- Q43. The nuclear g-factors of <sup>1</sup>H and <sup>14</sup>N are 5.6 and 0.40 respectively. If the magnetic field in an NMR spectrometer is set such that be protons resonates at 700 MHz, the <sup>14</sup>N nucleus would resonate at [NET June 2015]
  - (a) 1750 MHz
- (b) 700 MHz
- (c) 125 MHz
- (d) 50 MHz
- Q44. The <sup>1</sup>H NMR spectrum of a dilute solution of a mixture of acetone and dichloromethane in CDCl<sub>3</sub> exhibits two singlests of 1:1 intensity. Molar ratio of acetone to dichloromethane of thane in the solution is

[NET June 2015]

(a) 3:1

(b) 1:3

(c) 1:1

(d) 1:2

#### An organic compounds shows followings spectral data Q45.

[NET June 2015]

IR (cm<sup>-1</sup>) 1680

 $^{1}$ H NMR (CDCL,):  $\delta$ 7.66(m,1H), 7.60(m,1H), 7.10(m,1H), 2.50(s,3H)

 $^{13}$ C NMR (CDCl<sub>3</sub>);  $\delta$ 190,144.134,132,128.28m/z(EI):126(M+,100%),128(M+2,4.9%)

The structure of the compound is

The \*10 SnNMR chemical shift (approximately in ppm) corresponding Sn (relative to Me, Sn ) is Q46.

[NET June 2015]

(a) -4

(b) + 137

(c) + 346

(d) - 2200

Number of signals in the  ${}^{13}C\{{}^{1}H\}$  NMR spectrum of (R)  ${}^{4}$  methylpentan-2 - ol are Q47.

[NET June 2015]

(a) 3

(d) 4

(d) 6

The number of chemical shift non-equivalent protons expected in  ${}^{\dagger}H$  NMR spectrum of  $\alpha$  -pineneis Q48.

[NET Dec 2015]



α - pinene

(a) 7

(b) 8

(c) 9

(d) 10

The structure of the compoundS that matches the <sup>1</sup>H NMR data given below is Q49

NMR (DMSO -  $d_{ij}$ ):  $\delta$ 7.75 (dd. J = 8.8, 2.4 Hz. 1H). 7.58 (d. <math>J = 2.4 Hz. 1H), 6.70 (d. J = 8.8 Hz. 1H)

6.50 (broad s, 2H), 3.80 (s,3H)

[NET Dec 2015]

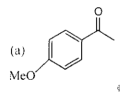
$$(a) \qquad NO_2 \qquad N$$

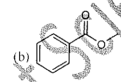
Q50. The  $^{1}$ H MNR frequency at 1.0 T is 42.4 MHz. If the gromagnetic ratios of  $^{1}$ H and  $^{13}$ C are  $27 \times 10^{-7}$  and  $6.75 \times 10^{-1}$  S respectively, what will be the  $^{13}$ C frequency at 1.0 T? [NET Dec.2015]

- (a) 10.6 MHz
- (b) 169.9 MHz
- (c) 42.6MHz
- (d) 21.3HMz
- Q51. In an NMR spectrometer containing a 2.5 T magnet, Larmor, Precession frequency of  ${}^{1}\text{H}$  is 100 MHz. The radiofrequency used in this spectrometer has an associated magnetic field strength of  $2.5 \times 10^{-4}\,\text{T}$ . The duration of a 90  ${}^{0}$  pulse in this instrument is [NET June 2016]
  - (a)  $25 \times 10^{-6}$  s
- (b)  $50 \times 10^{-6}$ s
- (c)  $25 \times 10^{-5}$  s
- (d) 50×10
- Q52. The correct structure of the compound, which shows following  $^{13}$ C NMR DEPT -135 data is  $^{13}$ C NMR DEPT-135 negative peaks at  $\delta$  30.2, 31.9, 61.8, 114.7 ppm; positive peak at 130.4 ppm [NET June 2016]
  - (a) OH
- (b) OF
- (c) 0
- (d) OH
- Q53. A compound displays the following spectral data. The correct structure of the compound is IR: 1690 cm
  - <sup>1</sup>H NMR:  $\delta$ 2.5(s,3H),3.8(s,3H),6.9(d,J=8Hz,2H), **7.8**(8.7=8Hz,2H)ppm

 $^{13}$ C NMR $\delta$ 197,165,130,129,114,56,26ppm

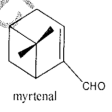
[NET June 2016]







Q54. In the  $^{1}H$  NMR spectrum of myrtenal, the two methyl groups are expected to display signals at (chemical shift values  $(\delta)$  in ppm) [GATE June 2016]



- (a) 1.35 (s, 3H) and 5.0 (s, 3H)
- (b) 0.74(s, 3H) and 1.33 (s, 3H)

(c) 1.22 (s, 6H)

- (d) 0.70 (s, 6H)
- Q55. The numbers of lines shown by the  $BH_3$  part of the molecule  $Ph_3P_3^{TI}BH_3$  in the  ${}^{4}H$  and  ${}^{41}B$  spectra are respectively  $\left[I({}^{11}B) = 3/2; I({}^{31}P) = 1/2\right]$  [NET June 2016]
  - (a) 8 and 8
- (b) 4 and 8
- (c) 3 and 6
- (d) 6 and 3

In a 200 MHzNMR spectrometer, a molecule, shows two doublets separated by 2 ppm .The observed coupling constant is 10 Hz. The separation between these two signals and the coupling constant in a 600 Q56. MHz spectrometer will be, respectively

(a) 600 Hz and 30Hz

(b) 1200 Hz and 30 Hz

(c) 600 Hz and 10 Hz

(d) 1200 Hz and 10 Hz

<sup>1</sup>H NMR spectrum of an organic compound recorded on a 500 MHz spectrometer showed a quartel with Q57. line positions at 1759, 1753, 1747, 1741Hz. Chemical shift  $(\delta)$  and coupling constant (Hz) of the quartet [NET Dec. 2016]

(a) 3.5 ppm, 6 Hz

(b) 3.5 ppm, 12 Hz

(c) 3.6 ppm, 6 Hz

(d) 3.6 ppm, 12 Hz

satellit**ės**, for including lines. spectral 19 F. NMR expected number of Q58. The INET Dec. 2016] A bundance  $\int_{-120}^{120} Xe(1 = 1/2) 26\%$ 

(a) two

(b) twenty one

(c)three

(d) one

H NMR spectrum of a mixture of benzene and acetonitrile shows singlets of equal integration. The molar Q59. [NET Dec.2016] ratio of benzene: acetonitrile is

(a) 1:1

(b) 2:1

(c) 1:2

[NET Dec. 2016]

The correct statements in the context of NMR spectroscopy is Q60.

(a) static magnetic field is used to induce transition between the spin states

(b) magnetization vector is perpendicular to the applied static magnetic field

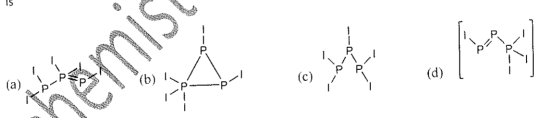
(c) The static magnetic fields is used to create population difference between the spin states

(d) static magnetic field induces spin-spin coupling.

The reaction between Pl., PSCL and zinc powder gives P, I, as one the products. The solution state 061  $^{31}P$  NMR spectrum of  $P_sI_s$  shows a doublet  $(\delta98)_s$  and a triplet  $(\delta98)_s$  . The correct structure of  $P_sI_s$ 

is

[NET Dec. 2016]



Number of lines in the  $^{19}\mathrm{F}$  NMR spectrum of  $^{12}\mathrm{F}$  (Br)-C(Br)Cl, at -120  $^{6}$  C assuming it a mixture of Q62@ [NET Dec. 2016] static conformations given below, are





(a) one

(b) two

(c) four

(d) five

Q63. The compound that exhibits following spectral data is

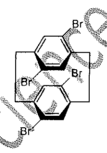
[NET Dec. 2016]

<sup>1</sup>H NMR:  $\delta 8.0(d, J = 12.3Hz, 1H), 7.7(d, J = 8.0Hz, 2H)$ 

6.8 (d, J = 8.0 Hz, 2 H), 5.8 (d, J = 12.3 Hz, 1H), 3.8(s, 3H), 3.0(s,6H) ppm

$$(a) \qquad \begin{matrix} N(CH_3)_2 \\ CO_2CH_3 \end{matrix}$$

Q64. Number of signals present in the proton decoupled <sup>13</sup>C NMR spectrum of the following compound is [NET Dec.2016]



(a) four

(b) six

(c) eight

(d) ten

Q65.  $^{13}$ C NMR spectrum of DMSO  $^{13}$ d, gives a signal at  $\delta$  39.7 ppm as a

[NET June 2017]

(a) singlet

(b) triplet

(c) quintet

(d) septet

Q66. The  ${}^{3}P\{{}^{1}H\}$  NMR spectrum of 2, 2, 6, 6 - N  ${}_{2}P_{4}CI_{4}$  (NMe) is expected to show

[NET June 2017]

(a) two triplets

(b) two doublets

(c) one doublet and one triplet

(d) one quartet and one doublet

Q67. The correct structure of the compound based on the following characteristic spectral data is **[NET Jun 2017]**IR: 1736 cm<sup>-1</sup>

 $^{1}\text{H NMR:}\delta3.59\big(s,3H\big),3.32\big(t,2H\big),2.25\big(t,2H\big),1.85-1.45\big(m,2H\big),1.73-1.62\big(m,2H\big)$ 

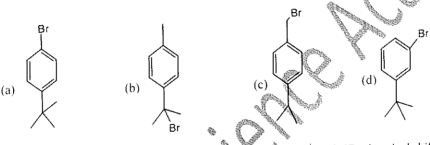
<sup>13</sup>C NMR:  $\delta$  174.0,51.0,32.9,32.9,32.8,31.0,23.0

$$(d) \qquad \stackrel{\mathsf{Br} \quad \mathsf{O}}{\longleftarrow}_{\mathsf{OMe}}$$

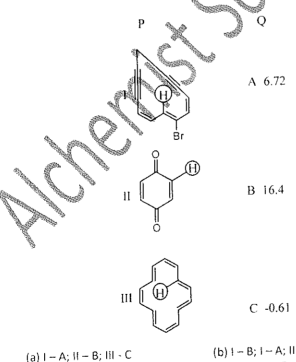
- For a certain magnetic field strength, a free proton spin transition occurs at 700MHz. Keeping the magnetic Q68. field strength constant the  $^{14}N$  -nucleus will resonate at  $\left(g\left(p\right)\approx5.6g$  and  $g\left(^{14}N\right)\approx0.4\right)$  [NET June 2017] (c) 200 MHz
- (b) 400 MHz The number of signals observed in the proton decoupled  $^{13}\mathrm{C~NMR}$  spectrum of the following compound (a) 700 MHz Q69. [NET Dec. 2017]



- (a) Five
- (b) Six
- (c) Ten
- (d) Thirteen
- The organic compound that displays following data is 'H NMR (400MHz) \*7.38(d) \*29(s)ppm Q70.



The correct match of the circled protons in Column P with  $^{1}H$  NMR chemical shift  $(\delta ppm)$  in Column Q is Q71. [NET Dec. 2017]



- (b) 1 B; 1 A; III C
- (c) I B; II C; III A
- (d) I C; II B; III A

- The g- factors of  ${}^{1}H_{-}$  and  ${}^{13}C_{-}$  are 5.6 and 1.4 respectively. For the same value of the magnetic fields Q72. strength. If the <sup>1</sup>H resonates at 600 MHz, the <sup>13</sup>C would resonate at [NET Dec. 2017]
  - (a) 2400 MHz (b) 600 MHz (c) 150 MHz (d) 38 MHz
- Q73. A compound shows following spectral data:

[NET June 2018]

<sup>1</sup>H NMR:  $\delta 7.9(d, J = 8Hz, 2H), 6.6(d, J = 8Hz, 2H)$ ,

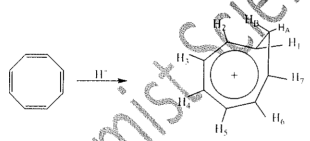
4.3 (q, J = 6.Hz, 2 H) 4.0 (br s, 2H, D<sub>3</sub>O exchangeable),

1.4 (t, J = 6 Hz, 3H)

Mass: m / z 165, 137, 120, 92

The correct structure of the compound is

The correct match of protons in Column-A with HI NMR chemical shifts in column-B for the product of Q74. the following reaction is [NET June 2018]



Column-A

Column-B  $(\delta ppm)$ 

(1) 0.3

(Q) H<sub>B</sub>

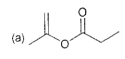
(ii) 5.1

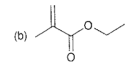
- (R) H<sub>1&7</sub>
- (III) 6.4
- (S)  $H_{2-6}$
- (iV) 8.5
- (a) P-H, Q-I, R-III,  $S\cdot IV$
- (b) P-1, Q-II, R-IV, S-III
- (c) P IV, Q I, R III, S III
- (d) P II, Q IV, R I, S III

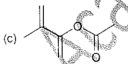
- In the  $^{31}P\{^{1}H\}$  NMR spectrum of a diamagnetic complex mer  $\left[M\left(PR_{3}\right)_{3}CI_{3}\right]$  ( M = transition metal, I = 0) Q75. [NET Dec. 2018] expected number of resonance(s) is (d) Six (c) Two (b) One
  - (a) Three

- The  $^{31}P\{^{1}H\}$  NMR spectrum of cis  $\left[Pt(PEt_{3})_{2}CI_{2}\right](^{198}Pt)$  (33.8% abundance) I =  $\frac{1}{2}$ ; its other isotopes are Q76. [NET Dec. 2018] NMR inactive;  ${}^{34}P$  ; I=1/2 ) is comprised with satellite peaks of a
  - (a) triplet
- (b) singlet
- (c) doublet
- (d) quartlet
- Structure of the compound displaying following characteristics spectral data. [NET Dec. 2018] Q77. IR: 1720 cm

 $^{1}H\ NMR: 6.2 (br, s1H), 5.5 (br, s. 1H) 4.2 (q, 2H), 2.0 (s, 3H), 1.1 (t, 3H)$  is



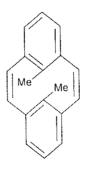




- Partial spectroscopic data is given below for an organic compound Q78.
- [NET Dec. 2018]
- (I) 4 signals between  $\delta$  120 150 ppm in  $^{13}$  C NMR spectrum
- (II) 2 doublets between & 6.8-8.5 ppm in <sup>1</sup>H NMR spectrum
- (III) an absorption band at 1724 cm in IR spectrum The structure of the compound is

ÒMe

The compound Pundergoes a pericyclic reaction under photochemical conditions to give compound Q. In Q79. compound Q. The relative stereochemistry and  $^{\dagger} H$  NMR chemical shift values of methyl groups (in  $\delta$ [NET Dec. 2018] ppm), respectively, are



- (a) cis; 5
- (b) trans; 17
- (c) cis; 17
- (d) trans; 5

Answer Key							
1. (a)	2. (b)	3. (b)	4. (c)	5. (a)	6. (d)	7. (d)	
8. (a)	9. (c)	10. (b)	11. (a)	12. (a)	13. (d)	14. (c)	
15. (b)	16. (a)	17. (d)	18. (a)	19. (d)	20. (a)	21. (d)	
22. (b)	23. (d)	24. (a)	25. (d)	26. (c)	27. (a)	28. (a)	
29. (a)	30. (a)	31. (b)	32. (d)	33. (c)	34. (a)	<b>35</b> . (a)	
36. (d)	37. (c)	38. (d)	39. (c)	40. (a)	41. (a)	42. (a)	
43. (d)	44. (b)	45. (d)	46. (d)	47. (d)	48. (d)	49. (b)	
50. (a)	51. (a)	52. (a)	53. (a)	54. (b)	55. <b>(a)</b>	56. (d)	
57. (a)	58. (c)	59. (c)	60. (b)	61. (c)	<b>6</b> 2. (c)	63. (c)	
64. (a)	65. (d)	66. (a)	67. (c)	68.(d)	69. (a)	70. (a)	
71. (b)	72. (c)	73. (a)	74. (a)	75. (c)	76. (b)	77. (b)	
78. (b)	79. (d)						

### **GATE Previous Year's Question**

The  $^{19}F$  NMR spectrum of  $PCI_{2}F_{\nu}\left(1\text{ for }^{31}P=1/2,1\text{ for }^{19}F=1/2\right)$  shows Q1.

[GATE 2000]

- (a) Two triplets and two doublets
- (b) Two triplet and one doublet
- (c) Two doublets and one triplet
- (d) three triplets and one doublet

The number of signals observed in  $^{4}$  H NMR  $^{2}$  spectrum of 3,5 - dibromotoluene is Q2.

[GATE 2001]

(a) 3

(b) 4

(c) 2

(d) 6

- In comparision to the frequency of the EPR transition, the NMR transition frequency is Q3.
  - (a) much higher
- (b) much lower
- (c) almost same

(d) none of these

F NMR spectrum of meriodional isomer of octahedral RhCl<sub>2</sub>F<sub>3</sub> complex, " Rh (I = 1/Q4.

[GATE 2001]

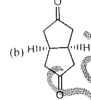
(a) one doublet

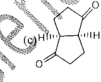
- (b) two doublets and one triplet
- (c) two doublets and two triplets

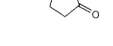
assuming  $J_{Rh-F} > J_{F-F}$  will show

- (d) one singlet and two triplets
- Among the bicyclo [3, 3, 0] octanediones given below, which one will exhibit FIVE signals in the broad band Q5. [GATE 2002] decoupled 13 C NMR spectrum

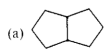








- The sensitivity of a 600 MHz NMR spectrometer is more than that of a 60 MHz spectrometer because Q6. [GATE 2003]
  - (a) Population of spin states is directly proportional to the applied magnetic field
  - (b) Population of spin states is inversely proportional to the applied magnetic field.
  - (c) According to the Bolzamann distribution law, the excess population in the lower spin state increases with increasing applied magnetic field.
  - (d) The spectral scan width is more for a 600 MHz spectrum compound to a 60 MHz spectrum.
- NMR spectrum of a compound with molecular formula  ${
  m C_4H_9NO_3}$  shows  $\delta$  5.30 (broad, 1H) , 4.10(q, Q7. 2H), 2,80 (d, 3H), 1.20 (t, 20 (t, 3H) ppm. The structures of the compound that is consistent with the above [GATE 2003] data is
  - (a) CHANHCOOCH, CH,
- (b) CH<sub>3</sub>CH<sub>2</sub>NHCOOCH<sub>3</sub>
- (c) CH,OCH,CONHCH,
- (d) CH<sub>3</sub>CH<sub>2</sub>OCH<sub>2</sub>CONH<sub>2</sub>
- Proton decoupled  $^{13}C$  NMR spectrum of a bicyclooctane  $\left(C_{s}H_{\mu}\right)$  exhibits only two signals. The structure Q8. [GATE 2004] of the compound is:









Q9. The vicinal coupling constant J expected for the protons  $H_p$  and  $H_q$  in the compound given below will be

in the range

[GATE 2004]

(a) 0 - 2 Hz

(b) 4 - 6Hz

(c) 8 - 10Hz

(d) 12 - 15<sub>\*</sub>H;

Q10. An organic compound having molecular formula  $C_6H_{11}BrO_2$  exhibits the following peaks in  $\frac{1}{2}H$  NMR spectrum. [GATE 2004]

 $\delta$  4.1 (2H, q, J = 7.5 Hz), 4.0 (2H, t, J = 7.5 Hz), 1.5 – 2.2 (4H, m), 1.25 (3H, t, J = 7.5 Hz)

The structure of the compound is:

$$(a) \bigvee_{\mathsf{Br}} \mathsf{O} \bigvee$$

(c)Br O

- (b) Br
- (d) Br
- Q11. 'H NMR spectrum of [18] annulene shows

[GATE 2005]

- (a) only one peak at  $\delta$  7.2 (18 Å)
- (b) Only one peak at  $\delta$  5.0 (18 H)
- (c) Two peaks at 8 9.0 (12 H) and 6 3.0 (6H)
- (d) two peaks at  $\delta$  9.0 (6H) and  $\delta$  3.0 (12 H)
- Q12. An organic compound having molecular formula  $C_x H_{12} O_2$  exhibits the following peaks in IR and  $^1H$  NMR spectra. [GATE 2005]

IR: 1720 (cm ) H NMR: 6.95(1H, d, J = 8.5Hz)5.90(1H, d, J = 8.5Hz, 4.53(1H,q,J = 6Hz)

1.41(3H, d, J = 6Hz), 1.20 (3H, s), 1.15 (3H, s)

(b) H<sub>3</sub>C

(c) H<sub>3</sub>C

- (d) CH<sub>3</sub>
  CH<sub>3</sub>
  CH<sub>3</sub>
- Q13. Which of the following compounds is expected to show a sharp singlet for one of its protons at

 $\delta \ge 8$  ppm in <sup>1</sup>H NMR spectrum, given that this single remains unaffected on shaking the solution thoroughly with D<sub>2</sub>O? [GATE 2006]

(a) CH,CO,H

(b) CH, CONH, CH,

(c)  $n - C_0 H_{13} C = CH$ 

(d)  $n - C_n H_{13}CHO$ 

- In the proton decoupled  $^{13}C$  and  $^{31}P$  NMR spectra of  $\left(CH_{3}\right)_{3}P=0$ , the number of lines observed, Q14. [GATE 2006] respectively, are

(a) two and one

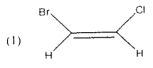
(b) one and two

(c) three and one

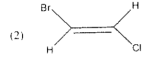
- (d) two and two
- Match the structures List-I with the coupling constant  $\left\lceil {}^{\dagger}HJ\left( Hz\right) \right\rceil$  given in List –II Q15.

List - !

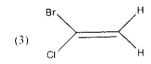
List - II



 $(i) \sim 1 \text{ Hz}$ 



 $(ii) \sim 10 Hz$ 



(a) 1-(i), 2-(ii), 3-(iii)

1-(ii), 2-(iii), 3-(i)

(c) 1-(iii), 2-(ii), 3-(i)

- (d) 1-(iii), 2-(i), 3-(ii)
- If CIF, were to be stereochemically rigid, its "FNMR spectrum  $\left[1 \text{ for } ^{16}\text{F} = \frac{1}{2}\right]$  would be (assume that CI Q16. [GATE 2008] is not NMR active)
  - (a) a doublet and a triplet
- (b) a singlet
- (c) a doublet and a singlet
- (d) two singlets

# Lined Answer Type for Q.17 and Q. 18

The reaction of  $_{
m PCI}$ , with methanol in the presence of triethylamine affords compound X. El mass spectrum of X shows a parent ion peak at m/z = 124. Microanalysis of X shows that it C, H, O and P. The  $^{\dagger}$ H NMR spectrum of X shows a boublet at 4.0 ppm. The separation between the two lines of the doublet is approximately  $15Hz \left( 1 \text{ for } ^{1}H \text{ and } ^{1}P = \frac{1}{2} \right)$ 

Q17. Compound X is [GATE 2008]

(a) (CH<sub>3</sub>O), P

- (b) (CH<sub>3</sub>O), P(O)
- (c) (CH<sub>3</sub>O), P(O)(OH)
- (d) (CH<sub>3</sub>O), PH

Q18. Upon heating, compound X is converted to Y, which has the same molecular formula as that of X. The 

'H NMR spectrum of Y shows two doublet centered at 3.0 ppm (separation of two lines ~20Hz) and 4.0 

ppm (reparation of two lines ~15 Hz) respectively.

[GATE 2008]

Compound Y is

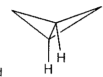
- (a)  $(CH_3O)_2 P(O)(OH)$
- (b) (CH<sub>3</sub>O), P(O)
- (c)  $(CH_3O)$ ,  $(CH_3)P(O)$
- (d) (CH<sub>2</sub>O)<sub>2</sub> (CH<sub>2</sub>)P(OH)

Linked Answer Q.19 and Q.20

[GATE 2008]

In the following reaction,

Q19. The reactive intermediate I and the product P are



(a) carbene and

(b) radical and



(d) radical and

Q20. The product P shows 'm' and 'n number of signals in <sup>1</sup>H and <sup>13</sup>C NMR spectra, respectively. The value of 'm' and 'n' are

(a) m = 3 and n = 3

(b) m = 2 and n = 3

(c)m = 2 and n = 2

(d) m = 4 and n = 4

Q21. The PNMR spectrum of P<sub>4</sub>S<sub>3</sub> Consists of

[GATE 2009]

(a) a singlet

- (b) a doublet and triplet
- (c) a doublet and quartet
- (d) two doublets

Q22. The <sup>1</sup>H NMR spectrum of HD consists of a

[GATE 2009]

(a) singlet

(b) 1:1 doublet

(c) 1:1:1 triplet

(d) 1:2:1 triplet

#### Q23. In the photochemical reaction

[GATE 2009]

MeO 
$$\frac{\text{hv}}{\text{CD}_3\text{OD}}$$
 [X]

Formation of the compound X can be inferred by the <sup>1</sup>H NMR signal at <sup>1</sup>H NMR spectrum of the strating material:

 $\delta$  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)  $\delta$  9.7 ppm
- (b)  $\delta$  7.8 ppm
- (c)  $\delta$  3.9 ppm
- (d) & 2.5 ppm

#### Common data for Q.24 and Q.25:

An organic compound  $X(C_{\nu}H_{\mu\nu}O)$  exhibited the following spectral data

IR: 1680 cm 1

'H NMR: 87.8(211.d.J7.5Hz), 7.2(2H, d, J = 75 z), 2.7 (3 H, s) and 2.4 (3H, s)

Compound X on treatment with m-choloroperbenzolc acid product two isomeric compound Y (major) and Z (minor)

## Q24. Compound Y and Z, respectively, are

[GATE 2009]

- Q25. Compounds Y and Z can be differentiated by carrying out basic hydrolysis, because
- [GATE 2009]

- (a) Y produces 4 -methylphenol and Z is unaffected.
- (b) Y produces 4 -methylphenol and Z produces 4-methyllbenzoic acid.
- (c) Y is unaffected and Z produces 4-methylbenzoic acid.
- (d) Y is unaffected and Z produces 4 methylphenol.

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

#### Common data for Q.26 and Q.27:

Treatment of  $W(CO)_6$  with 1 equivalent of  $Na(C_5H_5)$  is 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  $H_2$  produces P. (Use the 18 electron rule as your guide).

Q26. The compound M and N, respectively, are

- (a) [(C,H,)W(CO),]Na and [(C,H,)W(CO),H]
- (b)  $[(C,H_*)W(CO)_*]$  Na and  $[(C,H_*)W(CO)_*H]$
- (c)  $[(C_sH_s)W(CO)_s]$  Na and  $[(C_sH_s)W(CO)_sH]$
- (d)  $[(C_sH_s)W(CO)_s]$  Na and  $[(C_sH_s)W(CO)_sH]$
- Q27. The compound O and P, respectively, are

(a) 
$$[(C,H_c)W(CO)]$$
 and  $[(C,H_c)W(CO)]$  Br

(b) 
$$[(C,H_s)W(CO)_s]$$
 and  $[(C,H_s)W(CO)_sBr(TH_s)]$ 

(c) 
$$[(C,H,)W(CO),(THF)]$$
 and  $[(C,H,)W(CO)]$  Br

(d) 
$$[(C_5H_5)W(CO)_3]_2$$
 and  $[(C_5H_5)W(CO)_2]_2$  Br(THF)

#### Common data for Q. 28 and Q. 29:

An organic compound [X][C]H,O, exhibits the following spectral data

IR:~ 1720 cm

<sup>1</sup>H NMR 
$$2.35(s;6H)$$
,  $3.30(s,3H)$ ,  $3.83(t,2H)$ ,  $4.42(t,2H)$ ,  $7.07(s,1H)$ ,  $7.58(s,2H)$ 

The compound [X] with an excess of MeMgBr gives a 1 : 1 mixture of compounds [Y] and [Z]. The compound [Z] exhibits the following  ${}^{1}H$  NMR data

2.0 (bs, 1H) 3.30 (s, 3H), 3.56 (t, 2H), 3.70 (t, 2H)

Q28. The compound [X] is

[GATE 2010]

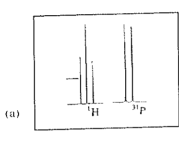
[GATE 2009]

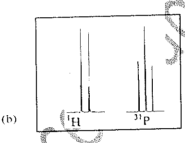
### Q29. The compound [Y] is

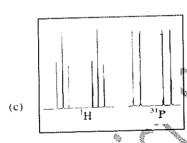
[GATE 2010]

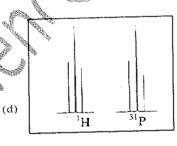
# Q30. The correct pair of ${}^{3}H$ and ${}^{31}P$ NMR spectral patterns for $C(H)(F)(PCl_{2})_{2}$

[GATE 2011]









# Statements for linked Answer Q 31 and Q.32

A ketone on treatment with bromine in methanol gives the corresponding monobromo compound [X] having molecular formula  $C_xH_xBrO_x$ . The compound [x] when treated with NaOMe in MeOH produces [Y] as the major product. The spectral data for compound [X] are:

'H NMR 81.17 (d.6H).3.02 (m.1H).4.10 (s.2H); CNMR817.37.39.210 .

# Q31. The compound [X]

[GATE 2011]

## Q32. The major product [Y] is

[GATE 2011]

Q33. In the proton decoupled <sup>13</sup>C NMR spectrum of 7 - norbornanone, the number of signals obtained is **[GATE 2012]** 

(a) 7

(b) 3

(c) 4

(d) 5

Q34. An organic compound Q exhibited the following spectral data:

[GATE 2013]

IR:

Q35.

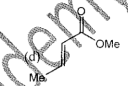
1760cm<sup>-1</sup>

 $^{+}H\ NMR: \delta(ppm); 7.2(IH,d,J=16.0Hz), 5.1(IH,m), 2.1(3H,s), 1.8(3H,d,J=7.0Hz) \\ 13CNMR: \delta(ppm:170) \\ (carbonyl\ carbon), \\ Compound\ Q\ is$ 

(a) O Me







Commond data for Q.35 and Q.36:

N, N -Demethylformamide (DMF) gives different patterns of signals for the methyl protons whenits 'H NMR spectrum is recorded at different temperatures.

Match the patterns of the NMR signals given in the Column with temperatures guves in the Column-II

[GATE 2013]

Column-II Column-II

- (i) Two singlets , for three protons each at  $\delta$  2.87 and 2.97 ppm
- (x) 25°C
- (ii) One sharp singlet for six protons at  $\delta 2.92$  ppm
- (y)  $120^{-0} C$

(iii) One broad single for six protons

(z)  $150^{\circ}C$ 

- (a) (i)-(x);(ii)-(y);-(lii)-(z)
- (b) (i) -(x);(ii)-(z);(iii)-(y)

- (c) (i) -(z);(ii) /(x)/ (iii) -(v)
- (d) (i)-(z);(ii)-(y);(iii)-(x)
- Q37. The number of signals that appear in the proton decoupled  $^{13}C$  NMR spectrum of benzonitrile (C.H.N)

[GATE 2013]

Q38. The set of protons (underline) in CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>OCH<sub>3</sub> that would exhibit different splitting patterns in high (500 MHz) and low (60 MHz) fields <sup>1</sup>H NMR , is [GATE 2014]

- (a) CH,CH,CH,OCH,
- (b) CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OCH<sub>3</sub>
- (c) CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>OCH<sub>3</sub>
- (d) CH<sub>3</sub>CH<sub>2</sub>CH<sub>3</sub>OCH<sub>3</sub>

Q39. The product of the following the following reaction gave 6 line  $^{13}$ C NMR spectrum with peaks at  $\delta$  175. 52, 50, 46, 37, 33 ppm. The structure of the product is **[GATE 2014]** 

- Q40 At room temperature, the number of singlet resonanes observed in the "H NMR spectrum of Me<sub>1</sub>CO<sub>1</sub>O<sub>1</sub>NMe<sub>2</sub> (NN dimethyl pivalamide) is \_\_\_\_\_\_ [GATE 2014]
- Q41. The beckmann rearrangement of a bromoatophenone oxime (C<sub>x</sub>H<sub>x</sub>BrNO) gives a major product having the following
  - "H NMR ( $\delta$  ppm): 9.89(s.IH). 7.88(s.IH). 7.45(d, IH). = 7.2Hz), 7.17 (m, 1H), 7.12 (d, 1H, J = 7.0Hz.), 2.06 (s, 3H). The structure of the product is. [GATE 2015]

Q42. The total number of lines expected (due to spin-spin coupling of proton with fluorine and deuterium nuclei) in the H NMR spectrum of the following compound is [GATE 2016]

Q43. The larmor frequency of  ${}^{1}H$  Teals (T) is 42.57 MHz. If the magnetogyric ratio for  ${}^{1}H$  and  ${}^{13}C$  and are  $26.75 \times 10^{-7}$  rad  ${}^{17}s^{-1}$  and  $6.72 \times 10^{-7}$  rad  ${}^{17}s^{-1}$ , respectively, the Larmor frequency of  ${}^{13}C$ , in MHz, at 1 Tesla will be \_[GATE 2016]

Q44. The structure of the compound having the following characteristics spectral data, is IR:  $1690 \text{ cm}^{-1}$   $^{1}\text{H NMR}: 1.30 (3\text{H. t. J} = 7.2\text{Hz}): 24\text{J} (2\text{H. q. J} = 7.2\text{Hz}): 2.32 (3\text{H. s}): 7.44 (1\text{H. t. -J} = 7.0\text{Hz}) 7.57 (1\text{H. dt J} = 7.0, 3.0 \text{ Hz}); 7.77 (1\text{H. t. J} = 3.0 \text{ Hz}); 7.90 (1 \text{ H. dt. J} = 7.0, 3.0 \text{ Hz}); El mass m/z 119 (100 %); 57 (80%)$ 

(a) 
$$CH_2CH_3$$
 (b)  $CH_2CH_3$  (c)  $CH_3$  (d)  $CH_2CH_3$ 

Q45. The  $^{^{19}}{\rm F~NMR}$  spectrum of CIF, when measured  $-60^{\,0}\,C$  will be observed as a

**IGATE 2016** 

(a) singlet

- (b) doublet
- (c) doublet and triplet
- (d) doublet of doublet and doublet of triplet

Q46. The <sup>13</sup>C NMR spectrum of acetone- d<sup>6</sup> has a signals at 30 ppm as a septet in the intensity ratio [GATE 2017]

- (a) 1:6:15:20:15:6:1
- (b) 1:3:6:7:6:3:1
- (c) 1:2:3:5:3:2:1

(d) 1 : 3 : 7 : 10 : 7 : 💃 🗓

Q47. The spectroscopic data for an organic compound with molecular formula  $C_{10}H_{12}O_2$  are given below. IR band around 1750 cm<sup>-1</sup>.

H NMR  $\delta$ 7.3(m:5H),5.85(q.1H, J = 7.2Hz),2.05(s.3H),1.5(d:3H, J = 7.2Hz)ppm. The compound is

- (a) methyl2-phenypropinate
- (b) 1-(phenylethyl) acetate
- (c) 2-(phenylethyl) acetate
- (d) methyl3-phenylpropionate

Q48. In the  $^{1}$ H NMR spectrum of an organic compound recorded on a 300 MHz instrument, a proton resonates as a quartet at 4.20 ppm. The individual signals of quartet appear a  $\delta$  4.17, 4.19, 44.21 and 4.23 ppm. The coupling constant J in Hz is [GATE 2018]

		Answer Key				
1. (a) 8. (d)	2. (a)	3. (b)	4. (*)	5. (d)	6. (c)	7. (a)
8. (d)	9. (c)	10. (c)	11. (c)	12. (c)	13. (d)	14. (a)
15. (c)	16. (a)	17. (a)	18. (c)	19. (a)	20. (c)	21. (c)
22. (c)	23. (a)	24. (b)	25. (b)	26. (a)	27. (a)	28. (b)
29. (b)	30. (c)	31. (d)	32. (c)	33. (b)	34. (a)	35. (b)
36. (30)	37. (5)	38. (b)	39. (c)	40. (3 to 3)	41. (a)	
42. (5.99 to 6.01)		43. (10.67 to 10.71)		44. (a)	45. (c)	46. (b)
47. (b)	48. (6)					

### **TIFR Pervious Year's Question**

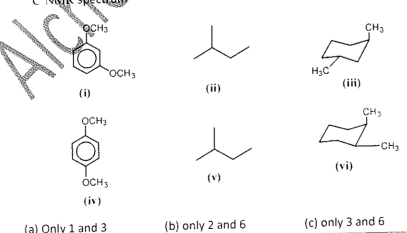
How many absorption lines will the following naturally occurring compound have in its proton-decupled Q1. [TIFR 2010] <sup>13</sup>C NMR spectrum

$$H_3C$$
 $H_3C$ 
 $CH_2CH_3$ 
(b) 4 (c) 5 (d) 6

(a)3For each of compounds below, choose the one in which the indicated hydrogen is farthest upfield in a Q2. [TIFR 2010] proton NMR spectrum:

(A) 
$$\frac{H}{1}$$
  $\frac{H}{2}$   $\frac{H}{3}$   $\frac{H}{3}$ 

At room temperature, which of the molecules are expected to give five NMR lines in the proton-decoupled Q3. [TIFR 2010] 13C NMR spectrum



(a) Only 1 and 3

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact: 011 -26511021, 8285787633, 8800927759, 9582285416

(d) All of the above

Q4. The <sup>13</sup>C NMR spectrum of a compound shows 6 peaks and the <sup>1</sup>H NMR spectrum shows 5 peaks. Which of the following is this compound [TIFR 2011]

(a) 
$$CH_1 - CH(CH_1) - CH(CH_2) - CH(CH_3) - CH_3$$

(b) 
$$CH_3 - C(CH_3), -CH_3 - CH_4 - CH_5$$

(c) 
$$CH_1 - CH(CH_1) - CH_2 - CH_3 - C(CH_3)_1 - CH_3$$

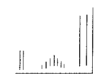
(d) 
$$CH_3 - CH(CH_3) - CH_2 - CH_2 - CH_3$$

Q5. Which of the following most closely resembles the <sup>13</sup> C NMR spectrum of ethanol Assume a scalar coupling of 150 Hz among the <sup>1</sup>H and the <sup>13</sup> C nuclei within a functional group, a scalar coupling of 50 Hz between the <sup>13</sup> C nuclei, a static magnetic field of 11.7 T and a temperature of 300K. [TIFR 2012]





Q6. Which of the following compounds would give the 'H NMR spectrum shown below [TIFR 2012]



- (a) CH,CH(CH,)CH,X
- (b) CH,C(CH,), CH,CH,X
- (c) CH,CH, (CH, )CH,X
- (d) CH,CH,CH(CH,)X
- Q7. The proton NMR spectrum of saturated hydrocarbon shows a single absorption line at 1.42 ppm with respect to TMS at room temperature. The area of the line is equivalent to 12 protons. Solely based o this observation, what are the tentative inferences you can draw about the nature of the hydrocarbon?

[TIFR 2012]

- (a) It is a pure compound i.e., there are no impurities present.
- (b) More than one conformation of the molecules may be present and they are undergoing rapid interconversion.
- (c) The hydrocarbon is cyclohexane.
- (d) All of the above
- Q8. The  $^{1}H$  NMR spectrum of a compound with molecular formula  $C_{3}H_{7}NO$  shows the following features:

Chemical shift (ppm)

6.50

2.25

1.10

[TIFR 2013]

Sphape

Broad singlet

quartet

triplet

Which of the following is in agreement with this information?

(a)  $(CH_1)$ , C = NOH

(b) CH, COCH, NH,

(c) CH,CH,CONH,

(d) HCO(NOH,),

Q9. The two fine-structure components of a nuclear magnetic resonance transition are observed at chemical shifts of 2.142 and 2.208ppm in a 300 MHz spectrometer Calculate the coupling constant [TIFR 2014]

(a) 19.8 Hz

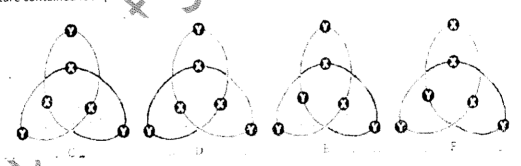
(b) 0.0666Hz

(c) 6.6 Hz

(d) data is insufficient

O10. Molecular knots have been observed in DNA and Proteins. Although synthetically challenging, a few purely organic molecular knots have also been reported in the literature. One such elusive knot is the trefoil knot. An organic trefoil knot was prepared by reacting two structurally distinct components X and Y. The reaction mixture contained four product as shown in the figure below

[TIFR 2015]



Which spectra will afford distinguishable spectral features for (1) C and D? And (2) E and F

- (a) Circular Dischroism Spectra for C and D;  $^1\mathrm{H}-\mathrm{NMR}$  for E and F
- (b)  $^{+}H-NMR$  and  $^{13}C-NMR$  spectra for C and D; Circular Dichroism for E and F
- (c)Absorbance and Emission Spectra for C and D; Circular Dichroims for E and F
- (d) Infra Red spectra for C and D; Circular Dichroism for E and F +

Q11.	Neopentyl chloride, $(CH_3)_3 CCH_2 Cl$ reacts with a strong base (sodium amide) to produce a new compound. This compound has two <sup>1</sup> H NMR singlets at $\delta 0.20$ ppm and $\delta 1.05$ ppm (intensity ratio = 2 : 3.					
	What is the Most probable structe	ure of this compound?	[TIFR 2016]			
	(a) 2 -methyl - 2 - butene	(b) I,I — dimethylcyclopropane				
	(c) methylcyclobutane	(d) cyclopentane				
Q12.	The <sup>1</sup> H NMR of I,I dibromoeth	nane consists of two well-separated signals	s, one large and another one			
	small. Which one of the following	statements is correct?	[THER 2016]			
	(a) The large signal is a quartet and	d the small single is doublet				
	(b) The large signal is a triplet and	the small signal is a singlet				
	(c)The large signal is a singlet and					
	(d) the large signal is a doublet and					
Q13.	A $C_sH_{12}O_2$ compound has strong	infared absorption at $3300$ to $3400$ cm $^{-1}$ .	The <sup>1</sup> H NMR spectrum has			
	three singlets $\delta 0.9, \delta 3.45$ and $\delta 3$	.2 ppm with relative areas 3 : 2 : 1 addi	tion of $D_2O$ to the sample			
	eliminates the lower field signal. Th	$10^{-13}$ C NMR spectrum shows three signals .	all at higher field than $\delta$ 100			
	ppm. Which of the following comp	oun <b>ds</b> best fits this data	[TIFR 2016]			
	(a) 1.5 -pentanediol	(b) 1,3 -dimethoxypropane				
	(c) 2.2 – dimethyl-13 -propanedi	(d) 2,4 -pentanediol				
Q14.	The <sup>1</sup> H NMR spectrum of a cor	npound A shows doublet and a septet. V	Which one of the following			
	statements is TRUE?		[TIFR 2016]			
Ā	(a) The spectrum is consistent with	A containing a CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> group				
<b>*</b>	(b) The spectrum is consistent with	A being $(CH_x)_2$ CHCI				

Q15. A compound of formula  $C_5H_{12}$  gives one signal in the  $^1H$  NMR and two singnals in the  $^{13}C$  NMR spectra. The compound is [TIFR 2016]

(c) The spectrum is consistent with A Containing a CH,CH, group

(d) The spectrum is consistent with A being  $(CH_2)_1 CCI_2$ 

(a) pentane

(b) 2-methylbutane

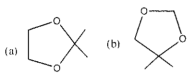
(c) 2.2 - dimethylpropane

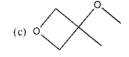
(d) cannot tell without more information

- Q16. Predict the multiplicities for hygrogens on CI, C3 and C4 of butanone associated with the spin-spin coupling in its <sup>1</sup>H NMR spectrum [TIFR 2017]
  - (a) H's on Cl: Singlet; H's on C3: Doublet H's on C4: Triplet
  - (b) H's on Cl: Singlet; H's on C3: Triplet; H's on C4: Quartet
  - (c) H's on Cl: singlett; H's on C3 Quartet; H's on C4 Triplet
  - (d) H's on Cl Triplet; H's on C3: Doublets H's on C4, Triplet
- Q17. An organic compound has the following spectroscopic properties: Mass spectrometry: m/z (very small), 87 and 43 are the largest ions; H NMR δ1.4 and 3.9 ppm (both singlets, intensity ratio 3 :2 <sup>13</sup>C NMR: δ108.64 and 25 ppm Infrared spectroscopy; several strong absorptions in the 1000 to 1300 cm <sup>1</sup> region

Which of the following the most likely formula of this compound?

[TIFR 2017]







Q18.

[TIFR 2017]

NMR spectroscopy can be used to assay for drug binding to certain protein targets. One of the primary objectives of any binding assay is the quantifaction of the free and the bound form of a drug molecule at a certain concentration of the protein target. Let us assume that the fully bound drug exhibits  $^{\mathsf{I}}$  H chemical shift of  $\delta_{\mathsf{A}}$  while that of the free formm resonates at  $\delta_{\mathsf{B}}$  for the same proton (see figure above). If the exchange timescale (i.e. proportional to  $k_{\mathsf{OFF}}$  as  $k_{\mathsf{ON}}$  is diffusion limited) between the free form of the drug and its bound form is in microseconds, which of the following statements cannot be true assuming that the drug is only 50 % bound with its protein target?

- (a) The NMR linewidth of the observed transition (s) will be different from the free form of the drug.
- (b) There will be two resonances obtained in the NMR spectrum Protein-Drug complex: one of the free form while other for the bound from.
- (c) We will see a single resonance at a position  $\delta_{
  m eff}$  which is in between  $\delta_{
  m A}$  and  $\delta_{
  m B}$  .
- (d) Varying the concentration of the drug molecule while observing the NMR signatures will provide an estimate of the binding constant.
- Q19. A compound with molecular formula  $C_3H_{12}O_2$ , has strong infrared absorption at 3300 to 3400  $cm^{-1}$ . The  $^1H$  NMR spectrum showed three singlets at  $\delta0.90, \delta3.45$  and  $\delta3.20$  ppm with relative areas 3:2:1. Addition of  $D_2O$  to the sample eliminates the lower field signal. The  $^{13}C$  NMR spectrum shows three singnals all higher than 100 ppm. Which of the following compounds best fits this data? [TIFR 2017] (a) 1,5-pentanediol (b) 1,3-demethoxypropane
  - (c) 2,2 -demethyl-1,3-propandiol
- (d) 2,4 pentanediol

Q20. An organic compound  $(C_9H_{10}O_3)$  exhibited the following spectral data: IR: 3400.1680cm  $^{++}$ H-NMR: $\delta$ 7.8 (1 H, doublrd, J = 8Hz), 6.5 (1 H singlet) , 5.8 (1 H, signment,  $D_2O$  exchangeable) 3.9 (3 H, singlet), 2.3 (3H, singlet). The compound is,

Q21. A bottle contains hydrogen molecule in gaseous state. The nuclear wavefunctoin of this hydrogen gas is given by

$$\frac{1}{\sqrt{2}}(\alpha(1)\beta(2) - \beta(1)\alpha(2))$$

What would be the NMR spectrum of this hydrogen gas, assuming that the wavefunction does not change during recording of the NMR spectrum?

- (a) A single NMR peak
- (b) Two NMR peaks, each is a doublet
- (c) Three NMR peaks, with relative intensities 1:2:1
- (d) No NMR lines
- Q22. Identify the organic molecule which contains 66.6% carbon, 11.1% hydrogen. In infra-red spectrum of the molecule, bands are observed at 2941-2857, 1715 and 1640 cm<sup>-1</sup>. In proton NMR, three signals appeared at (i) 7,52 (q, 2H), (ii) 7,88 (s, 3H), (iii) 8.93 (t, 3H) in ppm scale. [TIFR 2019]
  - (a) Ethyl ketone ®
- (b) 2-Butanol
- (c) 1 Butanladehyde
- (d) None of them
- Q23. In the broadband decoupled <sup>13</sup>C NMR spectrum, the number of signals appearing for the two pyrenediols

- (a) five and eight
- (b) eight and eight
- (c) eight and sixteen
- (d) five and ten

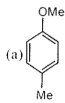
Answer Key						
1. (c)	2. (c)	3. (b)	4. (c)	5. (a)	6. (a)	7. (d)
8. (c)	9. (a)	10. (a)	11. (b)	12. (d)	13. (c)	14. (b)
15. (c)	16. (c)	17. (a)	18. (b)	19. (c)	20. (d)	21. (d)
22. (*)	23. (d)					

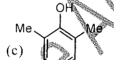
Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

#### Other examination previous Year's Question

- Q1. The chemical shifts of a doublet signal for a proton in a spectrum are 4.08 and 4.06 using a 400 MHz NMR spectrometer. The coupling constant (in Hz) is.
  - (a) 0.02
- (b) 8.0
- (c) 8.14
- (d) 10.0
- Q2. The <sup>1</sup>H NMR spectrum of 1,4-dimethoxybenzene will have
  - (a) ten singlets

- (b) two singlets
- (c) two doublets and one singlets
- (d) two doublets and two singlets
- Q3. The pattern of <sup>1</sup>H NMR spectrum of 14 dichlorobenzene is:
  - (a) AX
- (b) AM
- (c) AB
- Q4. An organic compound with molecular formula  $C_x H_{10}O$  exhibited 6 peaks in its broad-band decoupled  $^{13}C$  NMR spectrum. The possible structure of the compound is:



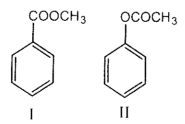


- Q5. The <sup>1</sup>H NMR spectrum of molecular has three distinct resonance at approximately, 3.4 ppm (a triplet with an intensity of 2 units), 1.6 ppm (a multiplet with intensity of 2 units). And at 1.0 ppm (a triplet with an units). The molecule could be
  - (a) (CH,CH,CH,), O

(b) CH,CH,CI

(c) CH,CH,CH,CH,

- (d) CH, PCI,
- Q6. The number of singles expected for the compounds m-and p-dichlorobenzenes in the broadband decoupled <sup>13</sup>C NMR spectra are, respectively,
  - (a) 6 and 4
- (b) 4 and 4
- (c) 4 and 2
- (d) 3 and 2
- Q7. In the  $^{1}$ H NMR spectra, signals due to methy groups in isomeric compounds I and II appear respectively at  $\delta$



- (a) 1.25 and 3.9
- (b) 3.9 and 2.1
- (c) 3.9 and 7.25
- (d) 7.25 and 2.1

- The low temperature  $^{19}$  F NMR  $^{19}$  spectrum of  $^{11}$ Fs  $^{12}$  molecule in solution should exhibit which of the following Q8. patterns? (Ignore any magnetic coupling effects to the iodine nucleus). (b) One doublet and one quintet (a) One singlet (d) One quartet and one triplet (c)One singlet and one quintet How many signals will be observed in the proton decoupled <sup>13</sup>C NMR for hexamethylbenzene? Q9. (d)4(c) 3(b) 2(a) 1 The H NMR spectrum of an organic compound of molecular formula C<sub>4</sub>H<sub>8</sub> exhibited only a singlet at Q10.  $\delta=1.9$  ppm. The compound is: (b) cis-2-butene (a) 1 - butane (d) cyclobutane (c) trans- 2- butene The strength of the coupling between germinal protons in the following molecules Q11. (a) increases as the size of the ring increases (b) Decreases as the size of the ring increases (c)Remains the same (d) No relation between the size of the ring and the coupling The expected spectral pattern in the proton coupled <sup>13</sup>C NMR spectrum of a mixture of equal weights of Q12. CHCl, and 13 CHCl, would be The increasing order of chemical shift of the indicated proton in the following series of compounds is Q13.
  - The  $^{+}H^{-}$  NMR  $^{-}$  spectrum of  $\left(\eta^{s}-C_{s}H_{s}\right)$  Fe recorded at room temperature has Q14.

(b) III < II < I

(a) One singlet

(a) 1 < 11 < 111

- (b) One multiplet
- (c) Two singlets

(c) | < | | | | | |

(d) Two multiplets

 $\{d\} | 1 < 11 < 1$ 

- Q15. An organic compound with molecular formula  $C_3H_6Cl_2$  exhibits only one singal in the  $^1H$  NMR spectrum. The compound is:
  - (a) 2,2 -dichloropropane
- (b) 1,2 dichloropropane
- (c) 1,3 -dichloropropane
- (d) 1,1 dichloropropane
- Q16. The <sup>1</sup>H NMR chemical shift of CH<sub>3</sub>F<sub>3</sub>CH<sub>3</sub>Cl<sub>3</sub>CH<sub>4</sub>Br and CH<sub>3</sub>I are
  - (a) 2.16.2.68,3.05,4.26
- (b) 4.26,3.05,2.16,2.68
- (c) 4.26,3.05,2.68,2.16
- (d) 2.16,3.05,2.68,4.26
- Q17. The fine structure and intensity ratios expected in the proton NMR spectrum of  $^{14}$  NH $_4^+$  ion (for  $^{14}$  N, 1 = 1) are
  - (a) singlet

(b) boublet, 1:1

(c) triplet, 1:1:1

- (d) triplet 1:2:1
- Q18. In the 'H NMR spectrum of toluene, the resonance due to CH, group is expected at
  - (a)  $\delta$  0.5
- (b)  $\delta$  1.25
- (0)  $\delta$  2,  $\delta$
- (d) δ 3.5
- Q19. For any NMR active nucleus, the magnitude of radiofrequency required for observing nuclear magnetic resonance phenomenon depends on
  - (a) Strength of the magnetic fields.
  - (b) Choice of the nucleus
  - (c) Both on magnetic fields strength and choice of the nucleus.
  - (d) The nuclear energy levels.
- 20. In the proton NMR spectrum, an organic compound exhibited the following spectral data

$$\delta$$
 7.2 (1H, dd,  $J = 8$  and 1.5 Hz), 6.8 (1H, d,  $J = 1.5$  Hz), 6.7 (1H, d,  $J = 8$  Hz), 4.9 (2H, s), 3.9 (3H, s),

3.85 (3H,s), 3.5 (1H, br s, exchangeable with D,O)

The compound among the choice given below is

- Q21. In the broadband decoupled  $^{13}$  Carbon NMR spectrum, the number of singals expected for bicycle [2.2.1]— heptanes is
  - (a) 2

(b) 3

(c) 5

- (d)7
- Q22. Which of the following dinitrochlorobenzens exhibits two singlets its <sup>1</sup>H NMR spectrum?

- (b)<sub>NO2</sub> NO<sub>2</sub>
- (c)<sub>NO2</sub> NO<sub>2</sub>
- NO<sub>2</sub> (d) NO<sub>2</sub> CI
- Q23. In the broad band decoupled <sup>13</sup>C NMR spectrum, the number of signals appear for (a) catechol, (b) resorcinol and (c) hydroquinone, respectively are
  - (a) six, four and two

- (b) six, six and four
- (c) three, four and four
- (d) three, four and two
- Q24. The  $^{31}P$  NMR spectrum of  $PF_4N(CH_3)_2$  at room temperature and low temperature (173 K) respectively shows (assume that N and H do not couple)
  - (a) triplet and quintet

- (b) quintet and triplet
- (c) quintet and triplet of triplets
- (d) triplet and triplet of triplets
- Q25. The order of chemical shift ( $\delta$  value) in the  $^1H$  NMR spectrum of crotonaldehyde is
  - (a) Olefinic > CHO > Me
- (b) CHO > Me > Olefinic
- (c) CHO > Olefinite Me
- (d) Olefinic > Me > CHO
- Q26. How many signals would you expect to see for 1,4 dinitrobenzene in its <sup>1</sup>H NMR spectra?
  - (a) One in its 'H NMR and two in its 13 C NMR spectra.
  - (b) Two in its <sup>1</sup>H NMR and three in its <sup>13</sup> C NMR spectra
  - (c)Two in its <sup>1</sup>H NMR and one its <sup>13</sup> C NMR spectra
  - (d) One in its <sup>1</sup>H NMR and one in its <sup>13</sup> C NMR spectra.
- Q27. Which of the following will occur farthest downfields?
  - (a) The hydrogens of benzens
- (b) The hydrogens of dimethyl ether
- (c) The hydrogens of ethene
- (d) The hydrogens of ethyne

		····			
Q28.	How can CH <sub>3</sub> C	$H_2 - C (= O) - OCH$ , a	nd $CH_3 - C(= O) - OCH$	<sub>2</sub> CH <sub>3</sub> be distinguished by <sup>1</sup> H NMR ?	
	(a) The signals f	or each compound will l	have different multiplicitie	es.	
	(b) Only CII, -	$C(=O) \sim OCH_2CH_3$ wi	ill have a singlet, a triplet,	and a quarted.	
	(c) Only CH <sub>3</sub> Ci	$H_2 - C(= O) - OCH_3 $ w	ill have a singlet, a triplet,	and a quartet.	
	(d) It is by the m	ultiplicity of the signals	appearing farthest, down	field	
Q29.	How many singr	oals will vinyl chloride ha	ive in its <sup>1</sup> H NMR spectro	um?	
	(a) 1	(b) 2	(c) 3	(d) 4	
Q30.	Which of the fol	lowing compounds shov	vs(s) three signals its ${}^{\perp}H$	NMR spectrum?	
	(a) 2 - chloro -2-	methylbutane	(b) 3 - chloropentane		
	(c) 2 - cholopent	ane	(d) (a) and (b)		
Q31.	How many signa	ls does 2,2,4 - trimethy	Ipentane have in its H	IMR spectrum?	
	(a) 2	(b) 3	(c) 4	(d) 5	
Q32.	Why does the sig	nal for the hydrogen bo	nded to oxygen in ethanol	appears as a triplet in pure ethanol an	d
		anol that contains a tra			
	(a) Spin exchange	cause spin decoupling	(b) Electron exchange	cause spin coupling	
	(c)Photon exchan	ge causespin decoupling	g (d) Chemical exchange	cause spin decouling	
Q33,		t 100.		group has a coupling constant of 7 H.	Z
	What is the coup	ling constant for the CF	I <sub>3</sub> triplet?		
	(a) 0	(b) 3.5	(c) 7	(d) 10.5	
Q34.	Which of the follo	wing is not a true staten	nent ?		
	(a) 'Clean " splitti	ng patterns require the	chemical shift difference	between peaks to be at least 10 times	
	the coupling const				
	(b) Coupling const	ants do not depend on t	he operating frequency of	f the NMR.	
	(c)There are more	hert/ ppm on a 360 - Mi	Hz NMR than on a 60 - MF	HZNMR	
	(d) Spectra taken a	t higher operating frequ	iency have higher resoluti	on.	

Alchemist Science Academy					
	$^{-1}$ H NMR spectra of $\left(\mathrm{CH_3}\right)_2\mathrm{O},\mathrm{CH_3}\mathrm{F}$ and RCOOH s	how chemical shift $(\delta$ $)$ in ppm at			
Q35.	H NMK spectra of $(CH_3)_2$ occurs, and the				
	(a) 3.27, 4.30 and 10.8 respectively	(b) 4.30, 3.27 and 10.8, respectively			
	(c)3.27, 10.8 and 4.30 respectively	(d) 10.8, 4.30 and 3.27, respectively			
-00	Many many different types of protons (given unique	chemical shift in <sup>1</sup> H MNR are present in styrene?			
Q36.	NOW Many difference of Page 2019	/ J) (			
	(a) 3 (b) 4	(c) 5			
Q37.	The low temperature <sup>1</sup> H NMR spectrum of the t	he Fischer carbene complex [(CO), Cr(C(COH,)Me)]			
	shows for the methoxy group				
	(a) Two singlet of inequal intensity				
	(b) Four resonances in the ratio 1:3:3:1 due to co	oupling with Me group			
	(c) One single resonance for the methyl group				
	(d) Three peaks due to different environments for				
Q38.	The multiplicity of the signal in the ${}^{\rm M}{\rm P}-{\rm NMR}$ s	As			
	(a) 211.Rier	uartet with equal intensity			
	(c) Septer with discident into the	eptet with equal intensities			
Q39.		e number of single appear for (a) catechol, (b) resorcing	)1		
	and (c) hydroquinone, respectively are				
	(a) six, four and two (b) s	ix, six and four			
	(c) three rout are rou	hree, four and two			
Q40,	Which technique can be used to readily distingu	ish between isophthalic and terephthalic acids?			
	(b) IC NMR spectroscopy	R spectrosocopy			
	(c) Mass spectroscopy (d)	JV-Vis spectroscopy			
O41	. Cis and trans cinnamic, acids can be most readil	y distinguished and identified by,			
Q41	(a) IR spectra	(b) UV- spectra			
	(c) Chemical shift of the olefinic hydrogens				
	(d) Coupling constant of the olefinic hydrogens				
	Head office: 28-A. Jia Sarai ,	Hauz Khas New Dethi-110016,	74		

			Answer Ke	у		
1. (b)	2. (b)	3. (d)	4. (a)	5. (a)	6. (c)	7. (b)
8. (b)	9. (b)	10. (d)	11. (a)	12. (a)	13. (a)	14. (a)
15. (a)	16. (c)	17. (c)	18. (b)	19. (c)	20. (d)	21. (b)
22. (a)	23. (d)	24. (c)	25. (c)	26. (a)	27. (a)	28. (d)
29. (c)	30. (d)	31. (b)	32. (c)	33. (c)	34. (c)	28. (d) 35. (a)
36.(d)	37. (b)	38. (c)	39. (d)	40.(a)	41 <sub>n</sub> (d)	

### **Molecular Spectroscopy**

## NET/JRF Year's Question

01	In IR spectrum of $[Co(CN)_s]$	H the Co-H str	etch is observed at 184	0cm <sup>-1</sup> . The Co-D stretch in
Q1,	$\left[\operatorname{Co}(\operatorname{CN}), \operatorname{D}\right]^{3}$ will appear at			[NET June 2011]
	(a) 1300 cm <sup>-1</sup> (b) 1	400 cm <sup>-1</sup>	(c)1500 cm	(d) 1600 cm <sup>-1</sup>
Q2.	The rotational constant of 14	$N_2$ is 2 cm <sup>-1</sup> . The	e wave number of inc	ident radiation in a Raman
	spectrometer is $20487~\mathrm{cm}^{-1}$ .	What is the wave nu	imber of first scattered st	[NET June 2011]
	(a) 20479 (b) 2	20475	(c) 20499	(d) 20495
Q3.	The ${f Q}$ band in the vibrational	spectrum of acetyle	ne is observed in the	[NET June 2011]
	(a) C-C stretching mode	(b) C-H	symmetric stretching mo	ode
	(c) Bending mode	383	antisymmetric stretching	
Q4.	The vibrational energy levels,	$\upsilon''=0$ and $\upsilon=1$ of	a diatomic molecule are	separated by 2143 cm <sup>-1</sup> . Its
	anharmonicity $(\omega_{ m e} { m X}_{ m e})$ is 14 cr	$\mathfrak{n}^{\perp}$ . The values of $\omega$	$\left(\operatorname{in}\operatorname{cm}^{-1}\right)$ and first over	tone $(cm^{-1})$ of this molecule
	are respectively			[NET Dec. 2011]
	(a) 2143 and 4286 (b)	2 <b>15</b> 7 and 4286	(c) 2157 and 4314	(d) $2171$ and $4258$
Q5.	The molecule that will show R	aman spectrum, but	not IR spectrum, among	the following is
QJ.				[NET Dec. 2017]
	(a) H <sub>2</sub> (b)	HCI (c) Br(	(d) CS,	
Q6.	For a diatomic molecule AB,	the energy for the ro	tational transition form ${ m J}$	J=0 to $J=1$ state is 3.9 cm <sup>-1</sup>
	For a diatomic molecule AB,			[NET June 2012]
			(c)11.7 cm <sup>-1</sup>	(d) 15.6 cm <sup>-3</sup>
	1 - 1 - ·	7.8 cm <sup>-1</sup>		
Q7.	For the vibrational Raman s	pectrum of a hom	onuclear diatomic mole	cule, the selection rule under
	harmonic approximation is			[NET June 2012]
	(a) $\Delta v = 0$ only (b) $\Delta v = \pm$	lonly (c) $\Delta$	$v = \pm 2$ only (d)	$\Delta \mathbf{v} = 0, \pm 1$

		<del></del>					
Q8.	In the vibrat	ional spectru	m of CO <sub>2</sub> the r	number of fun	damental vibra	ational mode	s common in both infrared
	and Raman	are					[NET Dec. 2012]
	(a) Three		(b) Two	(	(c) One	(	d) Zero
Q9.	The relative	population ir	two states wi	th energies 13	$_{_{1}}$ and $_{\mathrm{E}_{2}}$ sati	sfying Boltzm	ann distributuion is given
	by $n_1/n_2 = ($	3/2)exp[-(	$\left[E_1 - E_2\right) / k_B T$	. The relative	degeneracy g	$_{2}/g_{+}$ is	[NET Dec. 2012]
	(a) 2		(b) 2/3	(c) 3/2		(d) 3	
Q10.	The absorpti	on spectrum	of $O_2$ shows a	vibrational str	ucture that be	comes contin	dumat 56875 cm <sup>-1</sup> . At
	the continuu	ım, it dissocia	ites into one g	round state a	tom $\left(\mathrm{O}_{_{\mathrm{g}}}\right)$ ar	nd on <b>e e</b> xc <b>it</b> e	d state atom $\left( \mathrm{O}_{\mathrm{c}} \right)$ . The
	energy differ	ence betweer	n $\mathrm{O}_{\mathfrak{e}}$ and $\mathrm{O}_{\!g}$ is	15125 cm	. The di <b>sso</b> cia	ti <b>o</b> n ene <b>rgy</b> (	$(in cm^{-1})$ of ground state
	of $O_2$ is:						[NET Dec. 2012]
	(a) $\frac{56875}{15125}$		(b) $\frac{15125}{56875}$	gi antibar.	cr72000	(c	u) 41750
Q11.	The vibration	nal frequency	and anharmo	onicity consta	<b>nt</b> of an alkal	i halide are	$300~\mathrm{cm^{-1}}$ and $0.0025$
	respectively.	The position	(in cm <sup>-1</sup> ) of it	ts fundamenta	al mode and fir	rst overtone a	are respectively.
	•	·					[NET Dec. 2012]
	(a) 300,600		(b) 298.5,595	5.5	c) 301.5, 604.5	(d	) 290,580
Q12.	In the presen	ce of an exte	nal magnetic f	ield (normal Z	(eeman effect)	the transitio	$n^{-1}D_2 \rightarrow P_1$ splits into
							[NET Dec. 2012]
	(a) 9lines		(b) 8 lines	(c	r) 7lines	(d	) 6 lines
Q13.	The equilibriu	m population	n ratio $\left(n_j/n_i^{}\right)$	of a doubly	-degenerate e	nergy level (	$\left( E_{j}  ight)$ lying $2$ unit higher
	than a lower r	non-degenera	te energy leve	$I\left(E_{j}\right)$ assum	ing K <sub>B</sub> T=1 u	nit, will be	[NET June 2013]
	(a) 2e <sup>-</sup>	(b) 2e <sup>2</sup>	(c) e <sup>-2</sup>	(d	) e <sup>-3</sup>		
Q14.	The atomic ma	asses of fluori	ne and hydrog	en are 19.0 a	nd 1.0amu, re	spectively ( 1	amu = $1.67 \times 10^{-27}  \text{kg}$
	). The bond ler	ngth of HF i	s $2.0 \overset{\scriptscriptstyle{0}}{ m A}$ . The r	moment of ine	ertia of HF is		[NET Dec. 2013]
	(a) $3.2 \times 10^{-4}$	<sup>7</sup> kg m <sup>2</sup>		(b) 6.4×	10 <sup>-47</sup> kg m	2	
	(c) $9.6 \times 10^{-47}$	kg m²		(d) 4.8×	10 <sup>-47</sup> kg m	2	

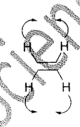
015		
Q15.	The $C=O$ bond length is 120 pm in $CO_2$ . The moment of the inertial	of CO <sub>2</sub> would be close to (masses
	of C and O are $1.9\times10^{-27}kg$ and $2.5\times10^{-27}kg$ , respectively)	[NET June 2014]
	(a) $1.8 \times 10^{-45} \mathrm{kg} \mathrm{m}^{-2}$ (b) $3.6 \times 10^{-45} \mathrm{kg} \mathrm{m}^{-2}$ (c) $5.4 \times 10^{-45} \mathrm{kg}$	g m <sup>2</sup> (d) $7.2 \times 10^{-45}$ k g m <sup>2</sup>
Q16.	Bond lengths of homonuclear diatomic molecules can be determined w	ith the help of both
	(a) Rotational and vibrational spectroscopy	[NET Dec. 2014]
	(b) Rotational and rotional Raman spectroscopy	
	(c) Rotational Raman and electronic spectroscopy	
	(d) Vibrational and electronic spectroscopy	
017	If the component of the orbital angular momentum along the molecu	lar axis of a heteronuclear diatomic
Q17.	If the component of the observe will show	[NET Dec. 2017]
	molecule is non-zero, the rotational-vibrational spectrum will show	
	(a) P and R branches only (b) P and Q branches only	
	(c) Q and R branches only (d) All the P,Q and R branches	
Q18.	If the bond length of a heteronuclear diatomic molecule is greater in	the upper vibrational state, the gap
Q10.	between the successive absorption lines of Paperen	[NET Dec. 2014]
	(a) Increase non-linearly	
	(c) Increase Linearly (d) Decrease linearly	
Q19.	intense band generally observed for a carbonyl group in the IR spectru	m is due to
	(a) The force constant of CO bond is large	[NET June 2015]
	(b) The force constant of CO bond is small	
	(b) The following the first the firs	
	(C) There is no change in dipole moment for CO bond stretching	
	(C) There is no change in dipole moment for CO bond stretching	
Q20.		[NET June 2015]
Q20.	(C) There is no change in dipole moment for CO bond stretching (d) The dipole moment change due to CObond stretching	[NET June 2015]
Ź	(C) There is no change in dipole moment for CO bond stretching  (d) The dipole moment change due to CObond stretching  The symmetry rotor among the following is  (a) CII, (b) CH <sub>3</sub> CI (c) CH <sub>2</sub> CI <sub>2</sub>	(q) C C 1 <sup>4</sup>
Q20.	(C) There is no change in dipole moment for CO bond stretching  (d) The dipole moment change due to CObond stretching  The symmetry rotor among the following is  (a) CH <sub>3</sub> (b) CH <sub>3</sub> Cl (c) CH <sub>2</sub> Cl <sub>2</sub> The spectroscopic technique, by which the ground state dissociation e	(q) C C 1 <sup>4</sup>
Ź	(C) There is no change in dipole moment for CO bond stretching  (d) The dipole moment change due to CObond stretching  The symmetry rotor among the following is  (a) CII, (b) CH <sub>3</sub> CI (c) CH <sub>2</sub> CI <sub>2</sub>	(d) $CCI_a$
Ź	(C) There is no change in dipole moment for CO bond stretching  (d) The dipole moment change due to CObond stretching  The symmetry rotor among the following is  (a) CH <sub>3</sub> (b) CH <sub>3</sub> Cl (c) CH <sub>2</sub> Cl <sub>2</sub> The spectroscopic technique, by which the ground state dissociation e	(d) CCl <sub>4</sub> nergies of diatomic molecules can be [NET June 2015]
Ź	(C) There is no change in dipole moment for CO bond stretching  (d) The dipole moment change due to CObond stretching  The symmetry rotor among the following is  (a) CH <sub>2</sub> (b) CH <sub>2</sub> Cl (c) CH <sub>2</sub> Cl <sub>2</sub> The spectroscopic technique, by which the ground state dissociation e estimated, is	(d) CCI <sub>4</sub> nergies of diatomic molecules can be  [NET June 2015] troscopy

Q22.	The molecule with	the smallest rotational	constant (in the microwave sp	pectrum) among the following is
				[NET Dec. 2015]
	(a) $N \equiv CH$	(b) HC ≡ CC	(c) $CCI \equiv CF$	(d) $B \equiv CCI$
Q23.	The spectroscopic	technique that can dist	inguish unambiguously betwee	en trans-1 , 2-dichloroethylene and
	cis-1, 2-dichloroetl	nylene without any num	erical calculate is	[NET Dec. 2015]
	(a) microwave spe	ctroscopy	(b) UV- visible spec	troscopy
	(c) X-ray photoelec	tron spectroscopy	(d) γ- ray spectros	сору
Q24.	If the reduced mas	s of a diatomic molecul	e is doubled without changing	its force constant, the vibrational
	frequency of the m	olecule will be	•	[NET Dec. 2015]
	(a) $\sqrt{2}$ times the	original frequency	(b) $\frac{1}{\sqrt{2}}$ times the	vibrational frequency
	(c) twice the original	al frequency	(d) unchanged	
Q25.	Upon application o	f the weak magnetic fi	el <mark>d, a line in the m</mark> icrowave a	bsorption spectrum of rigid rotor
	splits into 3 lines.	The quantum number	(#) of the rotational energy	level form which the transition
	originates is			[NET June 2016]
	(a) 0	b) 1	(c) 2	(d) 3
Q26.	The rotational cons	tant and the fundamen	tal vibrational frequency of H	Br are respectively, $10\mathrm{cm}^{-1}$ and
	2000 cm <sup>-1</sup> . The co	prresponding values for	DBr approximately are	[NET Dec. 2016]
	(a) $20 \text{ cm}^{-1}$ and $20 \text{ cm}^{-1}$	00 cm	(b) $10~\mathrm{cm}^{-1}$ and $14$	10 cm <sup>-1</sup>
	(c) 5 cm <sup>1</sup> and 20	00 cm	(d) $5~\mathrm{cm}^{-1}$ and $14$	10 cm <sup>-1</sup>
Q27.	Among the followin	g, both microwave and	rotational Raman active mole	cule is [NET Dec. 2016]
	(a) GH1	(b) N <sub>2</sub> O	(c) C ,H ,	(d) CO <sub>2</sub>
Q28.	Vibrations of diaton	nic molecules are usuall	y modelled by a harmonic pot	ential. If the potential is given by
	$x^2$ the correct state	ements is		[NET Dec. 2016]
	(a) force is 2x and fo	orce constant 2	(b) force is – 2x and force cor	nstant is 2
	(c) force is 2x and fo	rces constant is – 1	(d) force is – 2x and force cor	nstant is – 1

Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

- The v=0 to 1 vibration-rotation spectrum of a diatomic molecule exhibits transition for R(0),R(1),P(1)Q29. and P(2) lines at 2241,2254,2216 and  $2203~cm^{-1}$ , respectively. Form this data, we conclude that the [NET June 2017] molecule
  - (a) has rigid rotation and harmonic vibration
- (b) has anharmonic vibration
- (c) has rotational-vibrational interaction
- (d) is effected by nuclear spin-stastics
- The first electronic absorption band maximum of a polar and relatively rigid aromatic molecule appears at Q30. 310 nm but its flurescence maximum in acetonitrile solution appears with a large Stokes shift at 450 nm [NET June 2017] . The most likely reason for the Stokes shift is
  - (a) large change in molecular geometry in the excited state
  - (b) increase is dipole moment of the molecule in the excited state
  - (c) decrease in polarizability of the molecule in the excited state
  - (d) lowered interaction of the excited molecule with polar solvent
- The normal mode ethylene represented, by the figure below, is Q31.

[NET Dec. 2017]



(a) only IR active

- (b) only Raman active
- (c) both IR and Raman active
- (d) neither IR nor Raman active
- The pair contains a spherical top and a symmetric top, among the following, is [NET Dec. 2017] Q32.
- (b)  $Ch_3Cl_2.CH_3Cl$  (c)  $CH_3Cl.CH_4$  (d)  $CH_4.C(CH_3)$ ,
- Assuming harmonic approximation, the energy change for the reaction  $\, {
  m HCI+D}_{\,2} 
  ightarrow \, {
  m D\,CI+HD}\,$  in  $\, {
  m cm}^3$

(the vibrational frequency data in cm is given in the table below)

[NET June 2018]

HCI	D <sub>2</sub>	DCI	HD
2885	2990	1990	3627

(a) -258

(b) +258

(c) - 129

(d) +129

Q34. The transition moment integral for rotational transition between  $J=1:M_J=0$  and J=2:M=0 states for a diatomic molecule along the z-axis is proportional to [NET June 2018]

(a) 
$$\int_{0}^{\pi} \cos^{2}\theta \left(3\cos^{2}\theta - 1\right) d\theta$$

(b) 
$$\int_{0}^{\pi} \cos^{2}\theta \left(3\cos^{2}\theta - 1\right) \sin\theta d\theta$$

(c) 
$$\int_{0}^{\pi} \cos\theta \left(3\cos^2\theta - 1\right) \sin\theta d\theta$$

(d) 
$$\int_{0}^{\pi} \cos\theta (3\cos^2\theta - 1)\sin^2\theta\theta$$

Q35. A symmetric top molecule, among the following, is

(a) ethylene

(b) allene

(c) butatriene

(d) hexatriene

Q36. In the pure Raman rotational spectrum of  ${}^{16}O_2$ , whose electronic ground state is  ${}^{16}O_2$  transition to/from

(a) even J levels are missing

(b) odd J levels are missing  $\slashed{\#}$ 

(c) all J levels appear

(d) none of the J levels appear

Q37. In third and fourth lines in the rotational Raman spectrum of CO are separated by 8 cm<sup>-1</sup>. The CO bond length is given by [NET Dec. 2018]

(a) 
$$\sqrt{\frac{h}{16\pi^2\mu c}}$$

(b) 
$$\sqrt{\frac{3h}{32\pi^2\mu c}}$$

(c) 
$$\sqrt{\frac{h}{32\pi^2\mu c}}$$

(d) 
$$\sqrt{\frac{5h}{32\pi^2\mu c}}$$

Q38. A molecule AB, shows the following IR and IR Raman spectra

		.69	
$\overline{v}$ (cm <sup>-1</sup> )	IR	4	R <b>am</b> an 🌡
2215	Vs, PR		S) depol.
	-	CONCESSED OF	Selfer.
1250	Vs, PR		Vs, pol
	The same of the sa	46444	
560%	S, PQR		
4. 400			

The structure of the molecule is

[NET Dec. 2018]

(a) Linear symmetrical (D , ,

(b) Bent symmetrical  $(C_{2x})$ 

(c) Linear asymmetrical (2, )

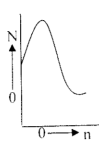
(d) Bent asymmetrical (C, )

#### **Answer Key**

			•			
1. (a)	2. (b)	3. (c)	4. (d)	5. (a)	6. (d)	7. (b)
8. (d)	9. (b)	10. (d)	11. (b)	12. (a)	13. (a)	14. (b)
15. (d)	16. (c)	17. (d)	18. (a)	19. (d)	20. (b)	21. (b)
22. (c)	23. (a)	24. (b)	25. (a)	26. (d)	27. (b)	28. (b)
29. (c)	30. (b)	31. (b)	32. (c)	33. (c)	34. (b)	35. (b)
36. (a)	37. (a)	38. (c)				

### **GATE Previous Year's Question**

Q1. The population (N) distribution over states (n) of a diatomic molecule corresponds to [GATE 200]



- (a) Translation
- (b) Vibration
- (c) Rotation
- (d) Electronic
  - [GATE 2002]

- Q2. Radiation of  $10^{14}$ Hz falls in the region of
  - (a) Radiofrequency
- (b) Microwave
- (c) Visible
- (d) X÷ray
- Q3. The spacing between the rotational lines of the HF is 40cm<sup>-1</sup>. The corresponding spacing between the rotational lines in DF is approximately [GATE 2002]
  - (a) 20 cm<sup>-1</sup>
- (b) 30 cm<sup>-1</sup>
- (c) 60 cm<sup>-1</sup>
- (d) 7.5 cm<sup>-1</sup>
- Q4. Neglecting the mass of hydrogen (1.0 amu) and deuterium (2.0 amu) with respect to that of iodine (127 amu), the ratio between fundamental vibrational frequencies HI and DI is [GATE 2004]
  - (a)  $\frac{1}{2}$
- (b) 2
- (c)  $\frac{1}{\sqrt{2}}$
- (d)  $\sqrt{2}$
- Q5. The population of  $J^{th}$  rotational level  $N_j$  is given by  $N_j = N_0 \left(2J + 1\right)e^{j(j+1)B_j + \Gamma}$ . The J value of rotational level with maximum population  $\left(J_{max}\right)$  is given by [GATE 2004]

(a) 
$$\frac{(2kT/B)-1}{\sqrt{2}}$$

$$\frac{\sqrt{2kT/B-1}}{2}$$

(c) 
$$\frac{kT}{B}$$

- (d)  $\frac{B}{kT}$
- Q6. Require matching of Items of column-I with the appropriate items in column-II. Choose the correct one form the alternate (a), (b), (c) and (d) [GATE 2005]

#### Column-

#### Column-II

#### Spectral Technique

#### **Selection Rule**

- P. Rotational transition
- I.  $\Delta v = \pm 1$
- o. Vibrational transition
- II.  $\Delta J = 0$
- R Electronic transition in atoms
- m.  $\Delta J = \pm 1$
- S. Molecular ensemble
- iv.  $\Delta l = \pm 1$
- $\mathbf{V} \cdot \Delta \mathbf{m}_1 = \pm 1$
- vi.  $\Delta v = 0$
- VII.  $\Delta l = 0$
- (a) P-I, Q-VI, R-VII, S-V
- (b) P-II, Q-I, R-IV, S-V
- (c) P-III, Q-I, R-IV, S-V
- (d) P-I, Q-VI, R-VII, S-V

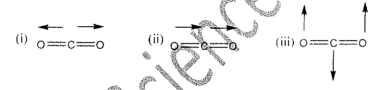
Q7.	The zero-point energ	y of the vibration of	35 Cl mimicking	oscillator with	a force constant
α,,,	$k = 2293.8 \text{ Nm}^{-1} \text{ is}$	y or one vibration of	C12 ////////////////////////////////////	oscinator with	[GATE 2006]
	(a) $10.5 \times 10^{-21}$ J	(b) 14.8×10 <sup>-21</sup> J	(c) 20×10 <sup>-21</sup> J	(d) 29.	6×10 <sup>-21</sup> J
Q8.	The selection rules for	the appearance, of P bra	nch in the rotational-	vibrational abso	rotion spectra of a
		in rigid rotor-harmonic os			[GATE 2007]
		_			
	(a) $\Delta v = \pm 1$ and $\Delta J =$	$\pm 1$ (b) $\triangle$	$\Delta { m V}=\pm$ 1 and $\Delta { m J}=$	+ 1	
	(c) $\Delta v = \pm 1$ and $\Delta J$	$=-1$ (d) $\Delta$	$\Delta { m V} = -1$ and $\Delta { m J}$	= -1	
Q9.	The $J=0 \rightarrow 1$ rotation	ial transition for ${}^{1}\mathrm{H}^{79}\mathrm{Br}$	occurs at 500.72G	Hz Assuming the	molecule to be a
	rigid rotor, the $J\!=\!3$ –				[GATE 2007]
	rigid fotol, the b 5	r transition occurs at			[0.712.2007]
	(a) 50.1 cm <sup>-1</sup>	(b) 66.8 cm <sup>-1</sup>	(c)16.7cm	(d) 83.5	5cm <sup>-1</sup>
Q10.	The rotational constant t	for $ m CO$ in the ground and	the first excited vibra	ational states are	$1.9$ and $1.6$ cm $^{\circ 1}$
	, respectively. The % cha	inge in the internuclear di	stance due to Vibratio	onal excitation is	
	(a) 9	(b) 30 (c) 16		(d) O	[GATE 2008]
	Linked Answer Type Q.1				
	The infrared spectrum	of a diatomic molecule	exhibits transition a	et 2144,4262	and $6354~\mathrm{cm}^{-1}$
	corresponding to excita	itions form the ground	state to the first, s	econd and, thir	d vibration states
	respectively				
Q11.	The fundamental transiti	on $(cm^{-1})$ of the diaton	nic molecule is at		[GATE 2008]
*	(a) 21 <b>57</b>	(b) 2170	(c) 2183	(d) 219	6
Q12.	The annarmonicity const	ant $\left(cm^{-1} ight)$ of the diator	nic molecule is		[GATE 2008]
*	(a) 0.018	(b) 0.012	(c) 0.006	(d) $0.00$	3
Q13.	The total number of way	ys in which two nonident	ical spin $-1/2$ part	icles can be orie	nted relative to a
•	constant magnetic field is		2) = port		[GATE 2008]
	(a) 1	(b) 2	(c) 3	(d) 4	

Q14.	1770	aman spectrum of	$^{19}\mathrm{F}_2$ shows a serie			
	19230.769 cm <sup>-1</sup> ,19	0227.238 cm <sup>-1</sup> and 192.	$23.707\mathrm{cm^{-1}}$ . The rotation	al constant for $^{19}F_2$ in $\mathrm{GHz}$		
	is:			[GATE 2009]		
		52 069	(c)105.936	(d) 3.531		
	(a) 26.484	(b) 52.968		(0)		
Q15.	The most populated ro	otational state for HCl(	$B = 8.5 \text{ cm}^{-1}$ ) at 300K is	[GATE 2010]		
	(a) 2	(b) 3	(c) 5	(d) 7		
Q16.	The ratio of life times	of two states that gives ri	se to line width of 1.0 cm <sup>-1</sup>	and 0.2 cm respectively is		
	(a) 1:2	(b) 1:5	(c) 2:1	(d) 5 : 1		
	Common Data for Q.1	17 and Q.18				
	A hypothetical molect	ule XY has the following p	roperties			
	Reduced mass: 2×10	<sup>-26</sup> kg X	-Y bond length: 100 pm	1		
	Force constant of the	bond: 8×10 <sup>2</sup> N.m <sup>-1</sup>				
Q17.	The frequency of rad	liation (in cm units) red	duired to vibrationally excite	e the molecule from $v=0$ to		
	v = 1 state is			[GATE 2011]		
	(a) 3184.8	(b) 2123.2	(c) 1061.6	(d) 840.0		
Q18.	The frequency of rad	iation (in cm <sup>-1</sup> units) requ	ired to rotationally excite th	be molecule from $J=0$ to $J=1$		
Q10.	state is			[GATE 2011]		
	(a) 1.4	(b) 2.8	(c) 3.2	(d) 3.6		
	Statement For Linked Answer for Q.19 and Q.20					
				O anly) to obtain the rotational		
	A 20491 cm 1 laser	line was used to excite o	sygen molecules (made of	O only) to obtain the rotational ecule has the first Stokes line at		
		e resulting rotational Kali	ian spectrum of oxygen mo.			
	20479 cm <sup>-1</sup>					
Q19.	The rotational consta	ant (usually denoted as B)	for the oxygen molecule is	[GATE 2012]		
	(a) 1.2 cm <sup>-1</sup>	(b) 2.0 cm <sup>-1</sup>	(c) 3.0 cm <sup>-1</sup>	(d) 6.0 cm <sup>-1</sup>		

Q20. The next rotational Stokes line is expected at

[GATE 2012]

- (a)  $20467 \text{ cm}^{-1}$  (b)  $20469 \text{ cm}^{-1}$  (c)  $20471 \text{ cm}^{-1}$  (d)  $20475 \text{ cm}^{-1}$
- Q21. The infrared spectrum of HCl gas shows an absorption band centered at 2885 cm<sup>-1</sup>. The zero point energy of HCl molecule under harmonic oscillator approximation is: [GATE 2013]
  - (a)  $2.8665 \times 10^{-22}$ J
- (b)  $2.8665 \times 10^{-20}$ J
- (c) 5.7330×10<sup>-2</sup> J
- (d)  $5.7330 \times 10^{-20}$  J
- Q23. Consider a two-state system at thermal equilibrium with equal degeneracy where the excited state is higher in energy than the ground state by  $0.1\,eV$ . The ratio of the population of the excited state to that of the ground state, at temperature for which  $k_BT=0.05eV$ , is [GATE 2016]
- Q24. Of the vibrational modes given below, the IR active mode(s) is(are) [GATE 2016]



- (a) (ii) only
- (b) (iii) only
- (c) (i) and (ii)
- (d) (ii) and (iii)
- Q25. The lowest energy of a quantum mechanical one-dimensional simple harmonic oscillator is 300 cm<sup>-1</sup>. The energy (in cm<sup>-1</sup>) of the next higher level is \_\_\_\_\_\_\_ [GATE 2017]
- Q26. For a diatomic vibrating rotor, in vibrational level v=3 and rotational level J, the sum of the rotational and vibrational energies is  $11493.6~\rm cm^{-1}$ . Its equilibrium oscillator frequency is  $2998.3~\rm cm^{-1}$ , anharmonicity constant is  $0.0124~\rm and$  rotational constant under rigid rotor approximation is  $9.7~\rm f^{-6}~cm^{-1}$ . The value of J is [GATE 2018]

(Upto nearest integer)

Q27. The spacing between the two adjacent lines of the microwave spectrum of  $H^{35}CI$ , is  $6.35\times1011Hz$ , given that bond length  $D^{35}CI$  is 5% greater than of  $H^{35}CI$  the corresponding spacing for  $D^{35}CI$ 

 $\times 10^{14} Hz$  . (Upto two decimal places)

[GATE 2018]

			Answer Key			
1. (c)	2. (c)	3. (a)	4. (d)	5. (b)	6. (c)	7. (b)
8. (c)	9. (b)	10. (a)	11. (b)	12. (c)	13. (c)	14. (a)
15. (b)	16. (b)	17. (c)	18. (b)	19. (a)	20. (b)	21. (b)
22. (38 to 42)		23. (0.13 to	0.14) 24.	. (d)	25. (899 to 901)	
26. (12) 27.	. (2.95)					

#### TIFR Previous Year's Question

- Q1. The stretching frequency of the O-H is about  $3600 \text{ cm}^{-1}$ . Compared to that, the stretching frequency of O-D and S-H bonds are very similar and about  $2500~{\rm cm}^{-1}$  . What can you conclude from these data [TIFR 2010]
  - (a) The electronic structure of O-D and O-H are same, and that S-H is different
  - (b) The force constant of the bond O-D and O-H is same.
  - (c) S-His a weaker bond than O-H or O-D bond
  - (d) All of these above
- Q2. Read the following two statements carefully.
  - 1. The changes in total angular momentum that occurs when a diatomic molecule (i. e. a rigid rotor) change rotational level from J=2 to J=3 is the same as the change in total angular momentum the occurs when an electron on a H atom changes from a d to an corbital, i. e. form l=2 to l=3
  - 2. The change in energy that occurs when a diatomic molecule (i. e. rigid rotor) changes rotational level from J=2 to J=3 is the same as the change in energy that occurs when a electron on a H atom changes from a d to an f-orbitals i. e. from I

Based on the above, which of the following is the correct statements

[TIFR 2010]

- (a) Both statement 1 and 2 are true
  - (b) Both statement 1 and 2 are false
- (c) statements1 is true, statement 2 is false (d) Statement 1 is false, and statement 2 is true
- The vibrational Raman effect, a considerably weak scattering phenomena, was first reported by Late Sir CV Q3. Raman in 1928. The intensity of the individual vibrational resonances observed in a Raman spectrum is [TIFR 2012] proportional to
  - (a) Number of molecules
- (b) Polarizibility of the bond
- (c) Wavelength of radiation used
- (d) All of the above
- Rotational energy diatomic molecules is given by  $E_{rot} = J(J+1)hB_e$  , where  $E_{rot}$  is in Joules. If the Q4. rotational constant for  $H_2$  molecule is given as  $B_c = 1.8324 \times 10^{12} Hz$ , the rotational period of the  $H_2$ molecule in J = 10 level will be [TIFR 2012]

- (a)  $1.33 \times 10^{-19}$  sec (b)  $5.0 \times 10^{-15}$  sec (c)  $5.46 \times 10^{-13}$  sec (d)  $7.39 \times 10^{-7}$  sec

Q5. The transition probability for spontaneous emission form state m to state n is given by an expression

$$A_{m \to n} = \left(\frac{64\pi^4 v_{mn}^3}{3hc^3}\right) \cdot \left(\left|\left\langle m\right| \hat{d} \mid n\right\rangle\right)$$

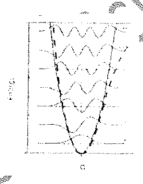
Where  $V_{\rm nm}$  is the frequency of transition, and the term in the parenthesis is the transition dipole. Assuming that magnitude of the transition dipole is same for all type of transitions, arrange the average lifetimes for the electronic, vibrational transitions in the proper order. [TIFR 2012]

- (a) electronic  $\leq$  vibrational  $\leq$  rotational
- (b) vibrational < rotational < electronic
- (c) rotational < vibrational < electronic
- (d) electronic < rotational ≈ vibrational
- Q6. Consider a classical harmonic oscillator with a mass 'm' and a force constant 'k' oscillating with a frequency
  - ${}^{\prime}v^{\prime}\,$  . Which of the following statements is NOT true for this system ?

[TIFR 2013]

- (a) 'v' increase if 'm' decrease
- (b) The oscillator is most likely to be found at its equilibrium position
- (c) The acceleration is maximum at its turning points
- (d)  ${}^{\prime}{}^{\prime}{}^{\prime}$  does not depend on how large the amplitude of the oscillation is
- Q7. Shown below in solid-line in the harmonic potential of a quantum oscillator for a diatomic molecule. If the harmonic potential is suddenly transformed into Morse potential shown in dashed-line how would the zero point energy shape of wavefunction change?

  [TIFR 2013]



- (a) Zero-point energy remains the same and wavefunction do not change
- (b) Zero-point energy remains the same but wavefunction reflects a change on the high Qside
- c) Zero-point energy changes and wavefunction reflects a change on the high Q side.
- Rotential never reflect any change in the shape of the wavefunction.
- Q8. Which of the following statements is/are true

[TIFR 2014]

- (i) HCl absorbs IR radiation
- (ii) CO<sub>2</sub> absorbs IR radiation
- (iii) H atom absorbs IR radiation
- (iv) H atoms UV -vis and microwave radiation

(a) (i) only

- (b) (i) and (ii) only
- (c) (i),(ii) and (iii) only
- (d) (i),(ii),(iii) and (iv)

Q9.	$\mathrm{N}_2$ does not shows pure vibrational spe	[TIFR 2014]	
	(a) triplet bond in $N_2$ is very strong	(b) The dipole moment of	$N_z$ is zero
	(c) Both (a) and (b)	(d) None of the above	
Q10.	Raman scattering is often seen overla	pping with fluorescence eman	ating form the sample. However
	fundamentally Raman process is differe	nt form fluorescence. This is bed	cause [TIFR 2014]
	(a) Raman scattering is a two – photon p	process and fluorescence is not	
	(b) Raman process is a scattering proces	s while fluorescence is not	
	(c) Raman process need not be stoke shi	fted	
	(d) Alf of the above		
Q11.	How many normal modes does the CO	, molecule have? What if the C	and the O atoms were constant to
	move in one dimension?		[TIFR 2015]
	(a) 4 normal modes for free ${\rm CO}_2$ and 4	for constrained $\mathbb{C}\mathrm{O}_2$	
	(b) 3 normal modes for free $CO_2$ and 2		
	(c) 3 normal modes for free CO <sub>2</sub> and 3	or constrained $\mathrm{CO}_2$	
	(d) 4 normal modes for free CO <sub>2</sub> and 2 i	for constrained $\mathrm{CO}_2$	
Q12.	Which of the following statements are tr	ue?	[TIFR 2015]
	(i) For a harmonic oscillattor potential, t	he spacing between adjacent e	nergy levels remain constant with
	increasing the quantum number.		
**	(ii) For a Morce oscillator potential, the	spacing between adjacent ener	gy levels increase with increasing
	the vibrational quantum number.		
	(fii) Harmonic oscillators are be used to ex	xplain the bond dissociation	
	(iv) Morse oscillator can be used to explain	n the vibration of molecule.	
	(a) i,ii and iii only(b) i and iv only	(c) i,ii and iV only (d	) i,ii,iii and iv only

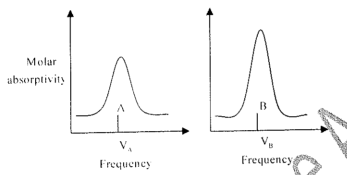
Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact : 011 -26511021, 8285787633, 8800927759, 9582285416

- For a harmonic oscillator in its ground state i. e. v=0 states, the energy is given by  $E=\frac{1}{2}hv$  where v Q13. [TIFR 2015] is the vibrational frequency. This is due to its

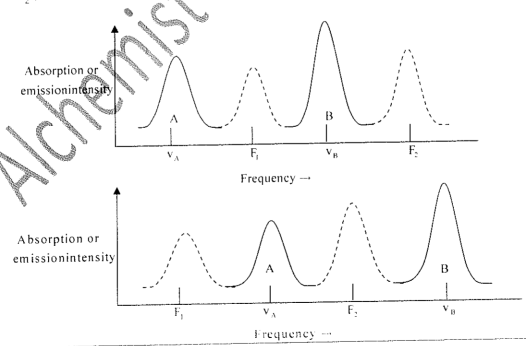
(a) Kinetic energy

- (b) Potential energy
- (c) Sum of kinetic and potential energy
- (d) Heat of formation
- The electronic absorption spectra or two species A and B are shown below Q14.

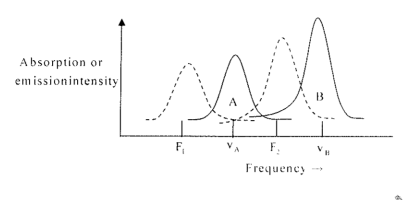
[TIFR 2015]

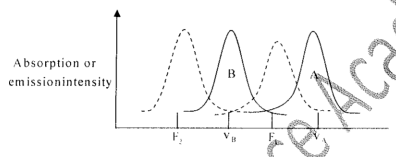


In a solution, these two species are dissolved, and they are moving freely. Using a tunable laser, when the solution is excited at  $V_{\Lambda}$ , a single fluorescence band is seen around frequency |F|. When the laser frequency is changed to  $|V_B|$ , two fluorescence bandare seen around frequencies  $|F_1|$  and  $|F_2|$ . Which of the following figures can qualitatively describe the correct relative position of the frequencies  $|v_{\rm A},v_{\rm B},F_{\rm I}|$  and  ${\rm F_2}$  ? The continuous line shows absorption and the dashed line shows fluorescence emission spectra.



Head office: 28-A. Jia Sarai , Hauz Khas New Delhi-110016 , Contact: 011 -26511021, 8285787633, 8800927759, 9582285416





Q15. Far infrared and microwave radiation is useful instudying the following process

[TIFR 2016]

- (a) Transition of inner electrons of atoms
- (b) Transition od outer (or valence) electrons in atoms or molecules
- (c) Changes in vibrational rotational states of molecules
- (d) Changes in molecular rotational only
- Q16. Isotopic substitution is an often-used procedure in assigning vibrational spectroscopic features but the difference in isotopic masses of the vibrating atoms has to be sufficiently large to make the vibrational shift observable. In a double isotopic replacement experiment, a proline with a  $^{13}C = ^{18}O$  carbonyl group is inserted in the middle of the polypeptide using molecular recombinant techniques. Given that the naturally found prolineth  $e^{12}C = ^{16}O$  group exhibits a narrow vibrational feature at  $11.711cm^{-1}$ , what is the shift of the carbonyl stretching vibration for the  $^{13}C = ^{18}O$  prolineisotomer.

**A**ssume:

- (A) The force constant for different isotopes can be consider identical.
- (B) The isotopic changes in the carbonyl group will have little effect on the vibrations of other atoms in proline
- (a) 80cm<sup>-1</sup>
- (b) 165cm<sup>-1</sup>
- (c) 85 c m<sup>-1</sup>
- (d) 42.5cm<sup>-1</sup>

- In a rotational microwave spectrum of  $\,\mathrm{C^{12}O^{36}}\,$  lines were equally spaced by  $\,3.663\,\mathrm{cm^{-1}}$  . In a rotational Q17. Raman spectrum of  $N_{\rm c}$  (normal isotope) the lines where equally spaced by  $8.04 {\rm cm}^{-1}$  . Assuming that the force constant for the two molecules is inversely proportional to their bond lengths, the ratio of the [TIFR 2017] vibrational frequency of CO to that of  $\,N_2\,$  will be
  - (a) 0.8368
- (b) 0.9952
- (c)1.0258
- (d) 1.2198
- Suppose you are carrying out an experiment measuring the Raman spectrum of  $N_2$  gas in the outdoor air. Q18. Where would you find a higher strength of the anti-Stokes line
  - (a) In Kanyakumari
  - (b) On top of Mt. Everest
  - (c) The strength will be the same in both the places
  - (d) Nitrogen would not have an anti-Stokes Raman line
- The fully symmetric C-H stretching mode  $\left(a_{1}\right)$  of  $CH_{1}$  was detected to be at  $3025 cm^{-1}$  . The C-HQ19. bending mode  $(t_2)$  on the other has was detected to be at  $1380 \, \mathrm{cm}^{-1}$  . If complete H/D exchange lebeling was done to produces the molecule  $CD_{\mu}$ , and the frequency ratio  $R_D$  is defined as  $= \left[ V_{\text{bend}} V_{\text{stretch}} \right] \text{ for } CD_4 R_H = \left[ V_{\text{bend}} / V_{\text{stretch}} \right] \text{ for ethane; which of the following statements is TRUE about}$  $R_{\rm H}?R_{\rm D}$  and the vibrational technique used for detection: [TIFR 2018]
  - (a)  $R_{\rm H}/R_{\rm D}=1.4$  while IR spectrum can be used to detect both the symmetric stretch and the bending mode
  - (b)  $R_{\rm H}/R_{\rm D}{=}1.0$  while Raman spectroscopy can be used to detect both the symmetric stretch and the bending mode
  - (c)  $R_{\rm H}/R_{\rm D}=1.0$  while IR can detect the symmetric stretch stretch and Raman the bending mode
  - (d)  $R_{\rm H}/R_{\rm H}$  while Raman can detect the symmetric stretch and IR the bending mode.
- The rotational constant for a diatomic molecule is  $1.9225 \, \mathrm{cm}^{-1}$  In general (Within the rigid rotor approximation), at T =  $600 {
  m K}$  , for a rotation state with maximum population  $\left({
  m J_{mix}}
  ight)$  and the position of maximum intensity of pure rotational absorption spectrum  $(I_{
  m max})$  , which of the following holds true

[TIFR 2018]

- (a)  $J_{\rm max}\!=\!7$  , while  $I_{\rm max}$  position is near transitions originating form J=7
- (b)  $J_{\rm max}\!=\!10$  while  $l_{\rm max}$  position is near transition originating from J=7
- (c)  $J_{max}\!=\!\!7$  , while  $I_{max}$  position cannot be determined from this information alone
- (d)  $I_{\text{max}} = 10$ , while  $I_{\text{max}}$  position cannot be determined form this information alone.

Answer Key						
1. (b)	2. (c)	3. (d)	4. (b)	5. (a)	6. (b)	7. (d)
8. (d)	9. (b)	10. (d)	11. (d)	12. (b)	13. (c)	14. (c)
15. (d)	16. (a)	17. (b)	18. (a)	19. (b)	20. (d)	

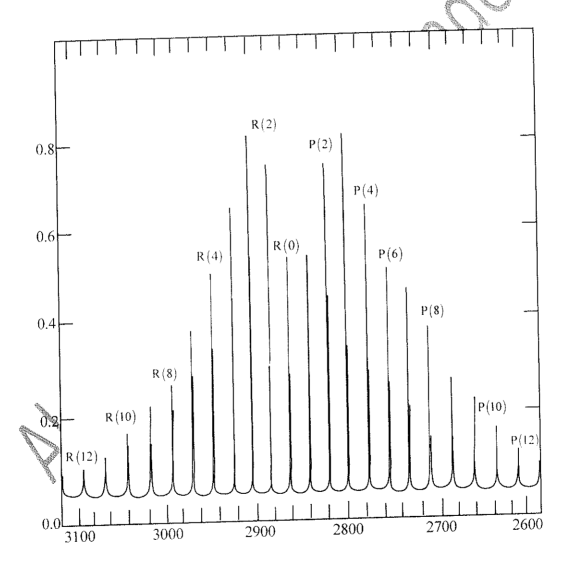
# Other Examination Previous Year's Question

- Q1. The lines in the microwave spectrum of  ${}^{1}H^{127}I$  are separated by A cm ${}^{1}$ . The lines in the microwave spectrum of  ${}^{2}H^{127}I$  will be separated by
  - (a) A/4
- (b) A/2
- (c)  $A/\sqrt{2}$

(d) A

## Common Information for Q.2 and Q.3.

Shown below is a vibrational – rotational spectrum for a diatomic molecule with reduced mass =  $10^{\circ}$  kg recorded at temperature T.



Q2. Moment of inertia  $(kgm^2)$  is:

- (a)  $1.0 \times 10^{-46}$
- (b)  $1.2 \times 10^{-47}$
- (c)  $5.0 \times 10^{-47}$
- (d)  $2.5 \times 10^{-47}$

Q3. Determine the correctness or otherwise Assertion  $\begin{bmatrix} a \end{bmatrix}$  and Reason  $\begin{bmatrix} r \end{bmatrix}$ .

Assertion [a]: Usually symmetric vibrations give rise to intense Raman lines. Non-symmetric ones are usually weak and sometimes not observed.

**Reason** [r]: Change I dipole is the highest during symmetric vibrations than the change during the non-symmetric ones.

- (a)Both  $\begin{bmatrix} a \end{bmatrix}$  and  $\begin{bmatrix} r \end{bmatrix}$  are the true  $\begin{bmatrix} r \end{bmatrix}$  is the correct reason for  $\begin{bmatrix} a \end{bmatrix}$
- (b)Both  $\begin{bmatrix} a \end{bmatrix}$  and  $\begin{bmatrix} r \end{bmatrix}$  are true but  $\begin{bmatrix} r \end{bmatrix}$  is not the correct reason for  $\begin{bmatrix} a \end{bmatrix}$
- (c) Both  $\begin{bmatrix} a \end{bmatrix}$  and  $\begin{bmatrix} r \end{bmatrix}$  are false
- (d)  $\begin{bmatrix} a \end{bmatrix}$  is true but  $\begin{bmatrix} r \end{bmatrix}$  is false

Q4. The force constant  $(in Nm^{-1})$  is:

- (a)  $2.9 \times 10^{-3}$
- (b) 5.8×10
- (c)  $1.2 \times 10^{\circ}$
- (d)  $5.8 \times 10^4$

Q5. Normal mode of vibration for the molecule  $C, H_{J}$  is

- (a) 18
- (b)15
- (c)12
- (d) 6

Q6. The vibrational energy of a diatomic may be written as (where Vis the frequency of vibration, and L is the moment of inertia of the molecules).

(a) 
$$(n+1/2)hv + \frac{h^2}{2I}J(J+1)$$
,  $n = 1,2,3...J$  and  $= 0,1,2....$ 

(b) 
$$(n+1/2)hv + \frac{h^2}{21}J(J+1), n = 0,1,2,3...$$
 and  $J = 0,1,2$ 

(c) 
$$(n+1/2)hv + \frac{\hbar^2}{21}M^2, n=1,2,3,...$$
 and  $M = 0,\pm 1,\pm 2,\pm 3,...$ 

(d) 
$$(n+1/2)hv + \frac{\hbar^2}{2I}M^2$$
,  $n = 0,1,2,3...$  and  $M = 0,1,2,3...$ 

Alchei	mist Science Acade	emy	CSITT				
Q7.	Given that the rotation	onal energy BJ (J+ 1	) , where the rotational const	ant has a value $\mathrm{B}=0.2\mathrm{cm}^{-1}$ and			
	kT is $209 cm^{-1}$ the approximate ratio of population of two rotational states with $J=10$ and $20$ are						
	(a) 1:2	(b) 2:1	(c) 19 : 27	(d) 21:41			
Q8.	The moment of inert	ia for HCl gas can be	determined form its microwav	e spectrum. Which property of the			
αυ.		obtained form the n		and the second second			
	(a) the vibrational fro	equency	(b) the force constant				
	(c) the bond strength	า	(d) the bond length				
Q9.	A wave of length $10$	nm is traveling at a	a speed of $10^6 \mathrm{m/s}$ . The frequ	ency <b>of</b> the wa <b>ve</b> is,			
	(a) $3 \times 10^{16} \text{Hz}$	(b) $10^{14} Hz$	(c) 3Hz	d) 3x [0]4Hz			
Q10.	The selection rules f	or the allowed rotation	on Raman lines is				
	(a) $\Delta J = 0, \pm 2$ (b)		(c) $\Delta J = 0$ (d) $\Delta J =$	$0,\pm1$ and $\pm2$			
Q11.	Replacement of hyd	Irogen by deuterium	atom bond to a heavy atom )	( in a polyatomic molecules would			
4	reduce the vibrational frequency of the X-Y stretching by a factor of						
	(a) 2	(b) $\sqrt{2}$		(d) 1.3			
Q12.	The first line in the r	otationāl Raman spe	tra of a diatomic molecules ap	pears with a stock shift of $12~{ m cm}^4$			
	. The Stoke shift for	W 34.					
	(a) 36 cm <sup>-1</sup>	(b) 24 cm <sup>-1</sup>	(c)18 cm <sup>-1</sup>	(d) 20 cm <sup>-1</sup>			
		a dibrational spectra	of AX exhibits a set of equall	y spaced lines with a separation of			
Q13.		ional constant of AX		, .			
dit.	(a) 10 cm	(b) 20 cm <sup>-1</sup>	(c) 5 cm <sup>-1</sup>	(d) 15 cm <sup>-1</sup>			
Q14.	45/ 1 May 1 May 1		al spectra of diatomic molecule	es when			
Q14.	(a) Anharmonicity is						
	(b) Anharmonicity is	s absent					
	(c) Vibrational and	rotational modes are	coupled				
	(d) An alternating e	lectric field is applied	,				
Q15.	The selection rule f	our observing rotatío	nal Raman spectrum is:				
	(a) $\Delta J = \pm 1$	(b) $\Delta J = \pm 2$	(c) $\Delta J = 0$	(d) $\Delta J = \pm 3$			

Q16.	The wavelength of light emitted when electron falls form the $\sqrt{n}=50$ orbits to the $\sqrt{n}=49$ orbits of F atoms, is						
	(a) 55nm	(b) 0.55cm	(c) $0.55 \stackrel{0}{\rm A}$	(d	) 55n		
Q17.	Which of the following n	nolecules has the	lowest vibrational stre	tching frequency?			
	(a) <sup>4</sup> H <sup>35</sup> C1	(b) <sup>2</sup> H <sup>35</sup> Cl	(c) <sup>4</sup> H <sup>36</sup> C1	(d	) <sup>4</sup> H <sup>37</sup> CI		
Q18.	A radiation which has ar	n energy of N 50	) kJ M ol <sup>-1</sup> falls in the	e following region	of the electromagnetic		
	spectrum.						
	(a) infrared	(b) visible	(c) ultraviol	et (d)	microwave		
Q19.	The microwave spectrum	n of a molecule yi	elds three rotational co	nstants. The mole	ecułe is		
	(a) Prolate symmetric top	o (b) Sph	nerical top				
	(c) Asymmetric top		(d) Oblate symmetric	<b>}</b>			
Q20.	A certain molecule can b	e treated as havi	ng only a doubly deger	nerate state lying	at 360 cm <sup>-1</sup> above the		
nondegenerate ground state. The approximate temperature $(K)$ at which $15\%$ of the molecules w							
	in the upper state is:						
	(a) 500	(b) 150	(c) 200		300		
Q21.	When $^{14}\mathrm{N}_2$ (with rotatio	- 723			ht, then the strokes and		
	anti-stokes lines for the n	n <b>ole</b> cüle is second	frotational state can b	e observed at			
	(a) 29412 cm and 292	410 cm <sup>-1</sup>	(b) 340.3 n	m and 339.9 n	m		
st.	(c) 14779.9 cm <sup>-1</sup> and 58	3529.9 cm <sup>-1</sup>	(d) 29384	(d) 29384 cm <sup>-1</sup> and 29424 cm <sup>-1</sup>			
Q22.	The magnitude of the nuc	clear spin angular	momentum of a nuclei	i $\sqrt{15}/2h$ units. T	he value of $f l$ is		
`	(a) 5/2	(b) 1/2	(c) 1	(d)	3/2		
Q23.	The rotational constant (	$B\big)$ of $H^{35}Cl,H^3$	<sup>2</sup> Cland D <sup>35</sup> Cl follow	s the order			
	(a) $H^{35}C1 > D^{35}C1 > H$	1 <sup>37</sup> C1	(b) $H^{35}Cl > H^{37}Cl$	> D <sup>35</sup> C1			
	(c) $D^{35}C1 > H^{35}C1 > H$	<sup>37</sup> CI	(d) $H^{37}CI > H^{38}CI >$	> D <sup>35</sup> Cl			

- At a given temperature, for a rigid rotor, the probability that the system is in the rotational state  $\,J=0\,{\rm is}\,$ Q24. 0.6 , in state J=1 is 0.3 , and 0.1 in J=2 . The energy in a rotational state J is given by  $J\left(J+1\right)B$ , where Bis the rotational constant. The average of the rotor at the give temperature is
  - (a) -6.0B
- (b) 1.2B
- (c) 3.6B
- (d) 4.8B
- The fundamental vibrational frequency Vof a homonuclear diatomic molecule with atomic Q25.

Mass m and force constant kis

- (a)  $v = \frac{1}{2\pi} \sqrt{\frac{k}{m}}$  (b)  $v = \frac{1}{v} \sqrt{\frac{2k}{m}}$  (c)  $v = \frac{1}{2\pi} \sqrt{\frac{k}{2m}}$

- An anharmonic diatomic oscillator has a frequency of  $3000\,\mathrm{cm}^{-1}$  and dimensionless anharmonicity of Q26. 1/30 . The diatoms is made to dissociate by supplying energy equal. To the potential well. The minimum relative kinetic energy of the ejected atom will be
  - (a) 1400 cm<sup>-1</sup>
- (b) 1450 cm<sup>-1</sup>
- (d) 1500 cm<sup>-1</sup>
- The moment of inertia of CO molecules is  $146\times10^{46} kg.m^2$  . The angular velocity in the J=1 energy Q27. level of CO molecules is
  - (a) 9.14×10<sup>11</sup> rad.sec<sup>-1</sup>
- (b) 10.20×10<sup>11</sup> rad.sec<sup>-1</sup>
- (c)  $11.44 \times 10^{11} \text{rad.sec}^{-1}$  (d)  $12.86 \times 10^{11} \text{rad.sec}^{-1}$
- The optimized variational wavefunction gives Q28.
  - (a) all properties and energy of same quality
- (b) properties better than the energy
- (c) energy better than properties (d) equal kinetic and potential energy values.
- For a simple diatomic molecules, the functional form of the potential energy curve can be expressed (with usual terms) as
  - (a)  $V(r) = D_e \left[ 1 + e^{-a(r+r_e)^2} \right]$
- (b)  $V(r) = D_e \left[ 1 e^{-a(r-r_e)^2} \right]$ 

  - (c)  $V(r) = -D_e \left[ e^{-a(r-r_e)^2} 1 \right]$  (d)  $V(r) = D_e \left[ 1 e^{a\left(1 e^{(r-r_e)^2}\right)} \right]$

- The mathematical form of transition dipole moment (with usual terms) for an optical transition form initial Q30. state  $\left(i\right)$  to final state  $\left(f\right)$  for visible radiation interaction with matter is given by
  - (a)  $\left\langle \psi_{f}\left|M\right|\psi_{i}\right\rangle$
- (b)  $\left\langle \psi_{\mathrm{f}} \left| \mu \right| \psi_{\mathrm{i}} \right\rangle$  (c)  $\left\langle \psi_{\mathrm{f}} \left| r.\mu \right| \psi_{\mathrm{i}} \right\rangle$  (d)  $\left\langle \psi_{\mathrm{i}} \left| \mu \right| \psi_{\mathrm{f}} \right\rangle$
- The bond length for  $H^9F$  is  $91.68\times10^{-12}m$  . Where does the axis of rotation intersect the molecular axes? Q31.
  - (a) Exactly in between F and H atoms
- (b) Axis of rotational is close to F
- (c) Axis of rotation is close to H
- (d) Far away to the right of F
- The number of IR active bands in  $H_2O$  molecules is Q32
  - (a) 2

- (b) 3
- (c)4

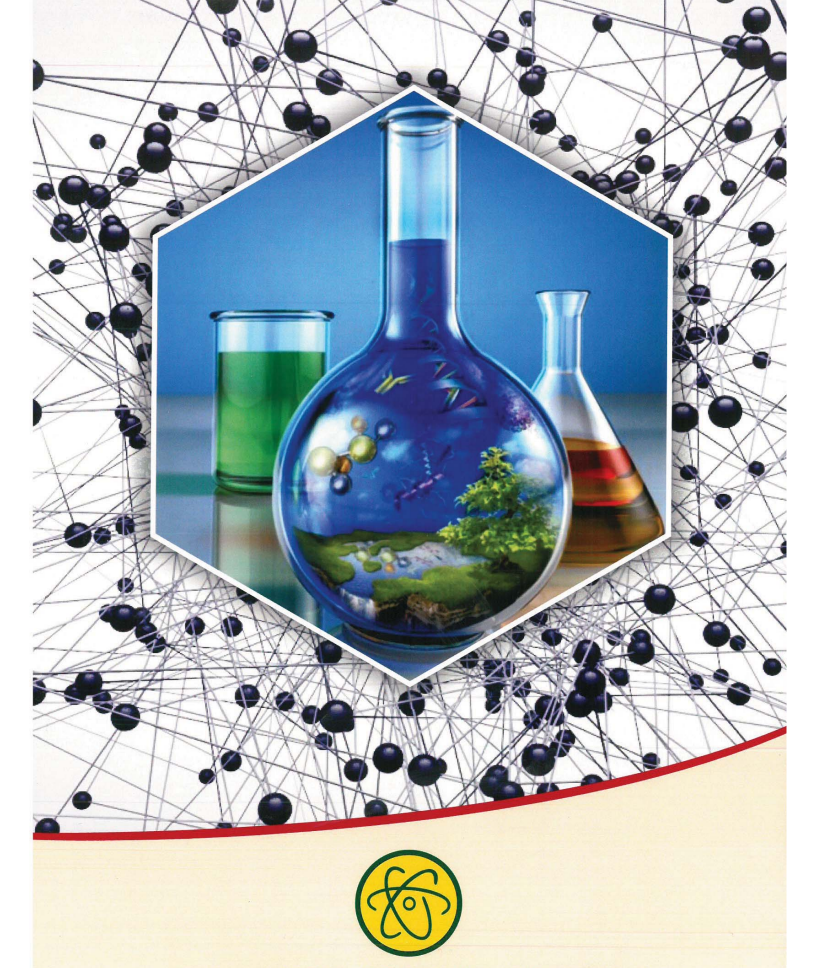
- The energy of rotational level with  $\,J=10\,$  , in wave number  $\left(cn^{2}\right)$ Q33.
  - (a) 2B
- (b) 110B
- (d) 10B
- The molecule which shows some common lines in the IR and Raman spectra is Q34.
  - (a) CO<sub>2</sub>

- (d) H<sub>2</sub>
- Which of the following molecules has a microwave spectra? Q35.
  - (a) CO,

- (d) OCS

- Which of the following molecules is infrared inactive? Q36.
  - (a) NO
- (c) H,
- (d) CH<sub>4</sub>

			Answer Key			
1. (b)	2. (b)	3. (*)	4. (a)	5. (c)	6. (c)	7. (d)
8. (d)	9. (b)	10. (a)	11. (b)	12. (d)	13. (c)	14. (a)
15. (b) 22. (d)	16. (b) 23. (b)	17. (c) 24. (b)	18. (a) 25. (b)	19. (c) 26. (c)	20. (c) 27. (b)	21.(d) 28. (a)
29. (b)	30. (b)	31. (b)	32. (b)	33. (b)	34. (c)	35. (d)
36. (c)						



# **ALCHEMIST SCIENCE ACADEMY**

28-/A, Jia Sarai, Hauz Khas, New Delhi-16 • Ph.: 011-26511021, 8285787633, 9582285416, 9149049297 www.csirnetalchemist.com • alchemistscienceacademy@gmail.com