

M.Sc.(Chem.) Semester - 4 (CBCS) Examination
April/May-2023 (NEW COURSE)
Modern Spectroscopy (Elective)

Time: 2:30 Hours

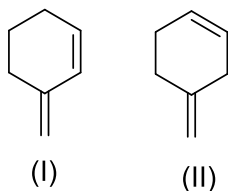
Marks: 70

Instructions:

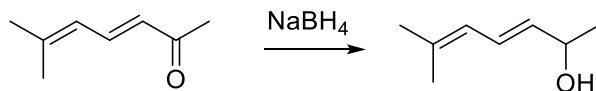
1. All questions are compulsory.
2. Figures to the right indicate marks.

Q.1(A) Answer the following questions. (04)

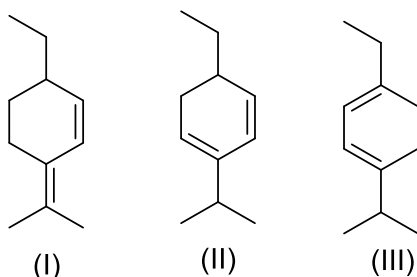
1. Give four structures of each (I) Auxochrome (II) Chromophore
2. How will you distinguish following pair of isomers using UV spectroscopy?

**Q.1(B) Answer the following question. (Any Two) (10)**

1. Discuss the effect of following factors on the shifting of $\lambda_{\max}$ with one example: (I) Exocyclic double bond (II) Ring residue
2. How can UV spectroscopy be used to determine the completion of the following reaction?



3. Arrange the following compounds based on their increasing value of $\lambda_{\max}$:

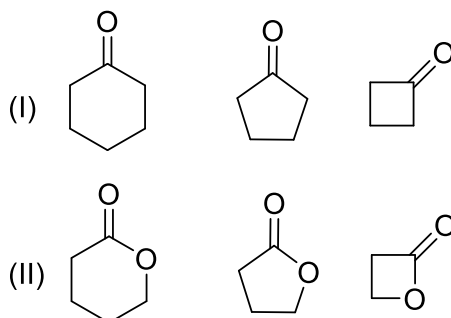
**Q.2(A) Answer the following questions. (04)**

1. Draw a schematic diagram of FTIR-spectrophotometer.
2. Why diatomic molecules Cl_2 and H_2 are IR inactive but HCl and H_2O are IR active?

Q.2(B) Answer the following question. (Any Two) (10)

1. Discuss various solid sample handling techniques in FT-IR and write its advantages.
2. Write a note on normal modes of molecular vibrations in the IR spectroscopy.

3. Arrange the following set of compounds (I & II) according to their increasing IR frequency. Give reasons for it.



Q.3(A) Answer the following questions.

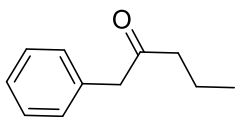
(04)

- How will you distinguish compounds having Cl and Br functional group using Mass Spectrum?
- Explain base peak and molecular ion peak for the benzyl alcohol.

Q.3(B) Answer the following question. (Any Two)

(10)

- Discuss FAB ionization method.
- Write a note on Time of Flight Mass analyzer including its advantages and disadvantages.
- The following compound shows peak at m/e 162, 134, 120 and 91 in the mass spectrum. Discuss its fragmentation for these peaks.



Q.4(A) Answer the following questions.

(04)

- Give difference between ^1H NMR and ^{13}C NMR.
- What is chemical shift? How it is represented? Give chemical shifts of most common solvents utilized in NMR.

Q.4(B) Answer the following question. (Any Two)

(10)

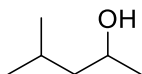
- Describe shielding-desielding effect of hybridization with suitable examples.
- A compound with Molecular Formula $\text{C}_7\text{H}_4\text{N}_2\text{O}_2$ showed Molecular Ion peak (M^+) at 148.12 m/z and base peak at 103.12 m/z . The FT-IR analysis showed characteristic bands at 1350 cm^{-1} , 1550 cm^{-1} , 2250 cm^{-1} . The PMR spectrum was obtained in the 400 MHz instrument using CDCl_3 as solvent and TMS as internal standard. The spectrum showed peak at δ ppm; 7.77 (t, $J = 8.8\text{ Hz}$, 1H), 8.02 (d, $J = 8.8\text{ Hz}$, 1H), 8.51 (d, $J = 8.8\text{ Hz}$, 1H), 8.57 (s, 1H). Analyze the given spectroscopic data, explain the signaling pattern and deduce the structure of unknown compound.

3. An unknown compound having Molecular Formula $C_{10}H_{12}O_3$ showed Molecular Ion peak (M^+) at 180.20 m/z. The FT-IR analysis showed characteristic bands at 1750 cm^{-1} . The PMR spectrum was obtained in the 500 MHz instrument using $CDCl_3$ as solvent and TMS as internal standard. The spectrum showed peak at δ ppm; 1.40 (t, 3H), 3.80 (s, 3H), 4.15 (q, 2H), 6.93 (d, $J = 8.0\text{ Hz}$, 2H), 8.01 (d, $J = 8.0\text{ Hz}$, 2H). ^{13}C NMR ($CDCl_3$): δ 15.20, 32.34, 64.23, 114.16, 122.94, 136.55, 162.63, 169.80. Analyze the given spectroscopic data, explain the signaling pattern and deduce the structure of unknown compound.

Q.5(A) Answer the following questions.

(04)

1. What is DEPT analysis? Discuss its application with one example.
2. Calculate the chemical shift of each carbon of following compound:



Q.5(B) Answer the following question. (Any Two)

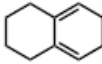
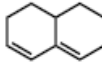
(10)

1. Write a note on 2D NMR spectrum and its applications.
2. An unknown compound having Molecular Formula C_9H_7N showed characteristic bands at 1450 cm^{-1} , 2235 cm^{-1} and 3100 cm^{-1} in the FT-IR spectrum. The PMR spectrum was obtained in the 400 MHz instrument using $CDCl_3$ as solvent and TMS as internal standard. The spectrum showed peak at δ ppm; 5.46 (d, $J = 10.8\text{ Hz}$, 1H), 5.88 (d, $J = 18.0\text{ Hz}$, 1H), 6.73 (dd, $J = 11.2\text{ Hz}$, 17.5 Hz , 1H), 7.49 (d, $J = 8.3\text{ Hz}$, 2H), 7.61 (d, $J = 8.3\text{ Hz}$, 2H). ^{13}C NMR ($CDCl_3$): δ 111.1, 117.7, 118.8, 126.7, 132.3, 135.3, 141.8. Analyze the given spectroscopic data, explain the signaling pattern and deduce the structure of unknown compound.
3. An unknown compound having Molecular Formula $C_5H_{10}O$ showed strong absorption bands at 1715 cm^{-1} in FT-IR. The PMR spectrum was obtained in the 100 MHz instrument using $CDCl_3$ as solvent and TMS as internal standard. The peaks appeared at δ ppm; 0.97 (d, 6H), 2.16-2.26(m, 1H), 2.3 (dd, 2H) and 9.85 (t, 1H). The ^{13}C NMR analysis showed peak at δ ppm: 24, 28, 54, 205. Analyze the given spectroscopic data, explain the signaling pattern and deduce the structure of unknown compound.

Standard Data Sheet for UV spectroscopy

UV Spectroscopy

Woodward Fieser Rules for Dienes

			
	<i>s-trans</i>	homoannular (cisoid)	heteroannular (transoid)
base values:	217 nm	253 nm	214 nm

Increments:

For each additional conjugated double bond	+ 30 nm
For each exocyclic double bond	+ 5 nm
For each alkyl group or Ring Residue	+ 5 nm

not affected by solvent

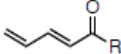
For each of the following groups:	
- OR	+ 6 nm
- O(C=O)R	+ 0 nm
- Cl	+ 6 nm
- Br	+ 6 nm
- SR	+30 nm
- NR ₂	+60 nm
- Ph	+60 nm

Where both types of cyclic dienes are present, the base with the longer λ_{max} is used.

UV Spectroscopy

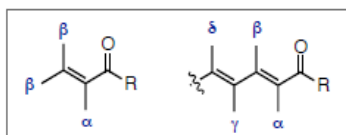
Woodward Fieser Rules for Enones

Solvent Correction	nm
H ₂ O	- 8
EtOH	0
CHCl ₃	+ 1
Dioxane	+ 5
Et ₂ O	+ 7
Hydrocarbon	- 11

				
	acyclic enone	6-membered ring enone	5-membered ring enone	acyclic dienone
base values:	215 nm	215 nm	202 nm	245 nm

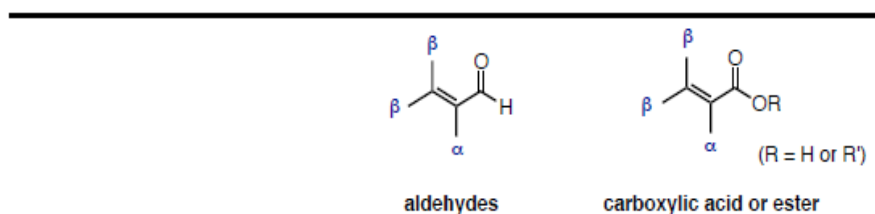
Increments:

	α	β	γ	δ and higher
For each additional conjugated double bond	+ 30 nm			
For each exocyclic double bond	+ 5 nm			
For each homodiene component	+ 39 nm			
For each alkyl group	+ 10 nm	+ 12 nm	+ 18 nm	+ 18 nm
For each of the following groups:				
- OH	+ 35 nm	+ 30 nm		+ 50 nm
- OR	+ 35 nm	+ 30 nm	+ 17 nm	+ 31 nm
- O(C=O)R	+ 6 nm	+ 6 nm		+ 6 nm
- Cl	+ 15 nm	+ 12 nm		
- Br	+ 25 nm	+ 30 nm		
- SR	+ 30 nm	+ 85 nm		
- NR ₂	+ 60 nm	+ 95 nm		



UV Spectroscopy

Woodward Fieser Rules for Other Conjugated Carbonyls



Base Values

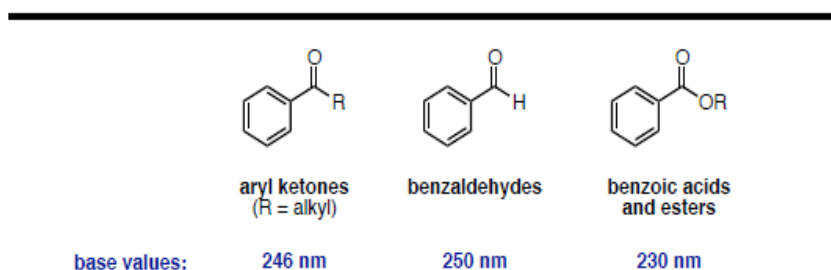
unsubstituted aldehyde or ester	208 nm	195 nm
with α or β alkyl groups	220 nm	208 nm
with α,β or β,β alkyl groups	230 nm	217 nm
with α,β,β alkyl groups	242 nm	225 nm

for an exocyclic α,β double bond	+ 5 nm
for an endocyclic α,β double bond in a 5- or 7-membered ring	+ 5 nm

as for enones, solvent correction is also relevant

UV Spectroscopy

Woodward Fieser Rules for Benzoyl Derivatives



Increments:

	ortho	meta	para
For each alkyl group or RR	+ 3 nm	+ 3 nm	+ 10 nm for RR +7 nm
For each OH or OR (R = alkyl)	+ 7 nm	+ 7 nm	+ 25 nm
For each O ⁻	+ 11 nm	+ 20 nm	+ 78 nm
For each of the following groups:			
- Cl	+ 0 nm	+ 0 nm	+ 10 nm
- Br	+ 2 nm	+ 2 nm	+ 15 nm
- NH ₂	+ 13 nm	+ 13 nm	+ 58 nm
- NH(C=O)CH ₃	+ 20 nm	+ 20 nm	+ 45 nm
- NHCH ₃			+ 73 nm
- N(CH ₃) ₂	+ 20 nm	+ 20 nm	+ 85 nm

Standard Data Sheet for IR spectroscopy

CHARACTERISTIC INFRARED ABSORPTION BANDS OF FUNCTIONAL GROUPS

Class of Compounds	Absorption, cm ⁻¹	Intensity	Assignment	Class of Compounds	Absorption, cm ⁻¹	Intensity	Assignment		
Alkanes and Alkyls	2850-3000	s	C-H stretch	Carboxylic Acids	2500-3500	s, broad	O-H stretch		
	1450-1470	s	C-H bend		R-C(O)-OH	1710-1715	s, broad	C=O stretch	
	1370-1390	m	CH ₃ C-H bend		C=C-C(O)-OH or Ar-C(O)-OH	1680-1710	s	C=O stretch	
	1365 + 1395 (two bands)	m	-CH(CH ₃) ₂ or -(CH ₃) ₃ bend	Esters	aliphatic 1160-1210	s-vs	O=C-O-C stretch		
	715-725	w	-(CH ₂) _n bend		acetates ~1240				
			aromatic 1250-1310						
Alkenes	3020-3140	w-m	=C-H stretch	R-C(O)-O-R	1735-1750	s	C=O stretch		
	1640-1670	vw-m	C=C stretch	C=C-C(O)-O-R or Ar-C(O)-O-R	1715-1730	s	C=O stretch		
	910 + 990 (two bands)	m + s	=C-H bend	R-C(O)-O-Ar	1760-1790	s	C=O stretch		
	RR'C=CH ₂	s	=C-H bend	Acyl Chlorides	R-C(O)-Cl	1785-1815	s	C=O stretch	
	cis-RCH=CHR'	m-s, broad	=C-H bend		Ar-C(O)-Cl	1770-1800	s	C=O stretch	
trans-RCH=CHR'	s	=C-H bend							
RCH=CR'R''	s	=C-H bend							
Alkynes	R-C≡C-H	3265-3335	s, sharp	≡C-H stretch	Anhydrides	R-C(O)-O-C(O)-R	~1750 + ~1815	s,s	C=O symmetric
		2100-2140	m	C≡C stretch		Ar-C(O)-O-C(O)-Ar	~1720 + ~1775	s,s	& asym. stretch
		610-700	s, broad	≡C-H bend			(both two bands)		
	R-C≡C-R'	2190-2260	vw-w	C≡C stretch	Nitriles	R-C≡N	2240-2260	m-s	C≡N stretch
				C=C-C≡N or Ar-C≡N		2220-2240	s	C≡N stretch	
Alkyl halides	R-F	1000-1350	vs	C-F stretch	Amines	R-NH ₂	~3400 + ~3500 (two bands)	w	N-H symmetric & asym. stretch
	R-Cl	750-850	s	C-Cl stretch			1580-1650	w-m	N-H bend
	R-Br	500-680	s	C-Br stretch		RR'N-H	3310-33350	w	N-H stretch
	R-I	200-500	s	C-I stretch					
Alcohols	3300-3400	s, broad	O-H stretch	Amides	R-C(O)-NH ₂	3200-3400 and 3400-3500 (two bands)	w-m	N-H symmetric & asym. stretch	
	C=C-CH ₂ -OH	1035-1050	m-s		C-O stretch		1650-1690	s, broad	C=O stretch
	R-CH ₂ -OH (1°) or C=C-CH(R)-OH	1050-1085	m-s		C-O stretch		1590-1655	m-s	N-H bend
	RR'CH-OH (2°) or C=C-CRR'-OH	1085-1125	m-s	C-O stretch	R-C(O)-NH-R	3400-3500	w-m	N-H stretch	
	RR'R''C-OH (3°)	1125-1205	m-s	C-O stretch		1640-1690	s, broad	C=O stretch	
	Ar-O-H	1180-1260	m-s	C-O stretch		1510-1560	m-s	N-H bend	
Ethers	R-O-R'	1085-1150	s	C-O-C stretch	R-C(O)-NR'R''	1630-1680	m-s	C=O stretch	
	Ar-O-R	1020-1075 and 1200-1275 (two band)	m-s	=C-O-C sym. & asym. stretch					
Aldehydes	2700-2725	m	H-C=O stretch	Nitro Compounds	R-NO ₂	~1550 and ~1370	s	N-O symmetric & asym. stretch	
	R-CH=O	1720-1740	s		C=O stretch	C=C-NO ₂ or Ar-NO ₂	~1525 and ~1335 (both two bands)	s	N-O symmetric & asym. stretch
	C=C-CH=O or Ar-CH=O	1685-1710	s	C=O stretch					
Ketones	RR'C=O	1710-1720	s	C=O stretch	Aromatic Compounds	3010-3100	m	Ar C-H stretch	
	C=C-C(O)-R	1665-1685	s	C=O stretch		1450-1600	m-s	ring C=C stretch	
	Ar-C(O)-R	1675-1695	s	C=O stretch		(two to four bands)	sharp		
	four member cyclic	1770-1780	s	C=O stretch		monosubstituted	730-770 and 690-710 (two bands)	s	C-H bend
	five member cyclic	1740-1755	s	C=O stretch					
	six member cyclic	1710-1720	s	C=O stretch		o-disubstituted	735-770	s	C-H bend
						m-disubstituted	750-810 and 690-710	s	C-H bend
						p-disubstituted	810-840	s	C-H bend
Intensity abbreviations: vw = very weak, w = weak, m = medium, s = strong, vs = very strong									

Intensity abbreviations: vw = very weak, w = weak, m = medium, s = strong, vs = very strong

Standard chemical shift value (ppm from TMS)

¹³C chemical shifts for some linear and branched-chain alkanes

Alkane	C-1	C-2	C-3	C-4	C-5
Methane	-2.5				
Ethane	5.7				
Propane	15.8	16.3	15.8		
Butane	13.4	25.2	25.2	13.4	
Pentane	13.9	22.8	34.7	22.8	13.9
Hexane	14.1	23.1	32.2	32.2	23.1
Heptane	14.1	23.2	32.6	29.7	32.6
Octane	14.2	23.2	32.6	29.9	29.9
Isobutene	24.5	25.4			
Isopentane	22.2	31.1	32.0	11.7	
Neopentane	31.7	28.1			
2,3-Dimethylbutane	19.5	34.3			
2,2,3-Trimethylbutane	27.4	33.1	38.3	16.1	
3-Methylpentane	11.5	29.5	36.9	(18.8, 3-CH ₃)	

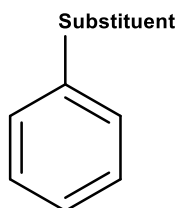
Olefinic carbons	
Substituent carbon C	Correction
α	+10.6
β	+7.2
γ	-1.5
α'	-7.9
B'	-1.8
γ'	-1.5
z (cis)	-1.1

Increments* (ppm) for substituents Y on alkanes

Substituent Y	α		β		γ
	Terminal	Internal	Terminal	Internal	
CH ₃	+9	+6	+10	+8	-2
HC=CH ₂	+20		+6		-0.5
C $\equiv$ CH	+4.5		+5.5	+2	-3.5
COOH	+21	+16	+3	+2	-2
COO ⁻	+25	+20	+5	+3	-2
COOR	+20	+17	+3	+2	-2
COCl	+33	+28		+2	
CONH ₂	+22		+2.5		-0.5
COR	+30	+24	+1	+1	-2
CHO	+31		0		-2
Phenyl	+23	+17	+9	+7	-2
OH	+48	+41	+10	+8	-5
OR	+58	+51	+8	+5	-4
OCOR	+51	+45	+6	+5	-3
NH ₂	+29	+24	+11	+10	-5
NH ₃ ⁺	+26	+24	+8	+6	-5
NHR	+37	+31	+8	+6	-4
NR ₂	+42		+6		-3
NH ₃ ⁺	+31		+5		-7
NO ₂	+63	+57	+4	+4	
CN	+4	+1	+3	+3	-3
SH	+11	+11	+12	+11	-4
SR	+20		+7		-3
F	+68	+63	+9	+6	-4
Cl	+31	+32	+11	+10	-4
Br	+20	+25	+11	+10	-3
I	-6	+4	+11	+12	-1

Data Sheet

Increments in the shifts of the aromatic carbon atoms of monosubstituted benzenes (ppm from benzene at 128.5 ppm)



Substituent	C-1	C-2 (<i>ortho</i>)	C-3 (<i>meta</i>)	C-4 (<i>para</i>)
H	0.0	0.0	0.0	0.0
CH ₃	+9.3	+0.7	-0.1	-2.9
CH ₃ CH ₂	+15.6	-0.5	0.0	-2.6
CH(CH ₃) ₂	+20.1	-2.0	0.0	-2.5
HC=CH ₂	+9.1	-2.4	+0.2	-0.5
C ₆ H ₅	+12.1	-1.8	-0.1	-1.6
F	+35.1	-14.3	+0.9	-4.5
Cl	+6.4	+0.2	+1.0	-2.0
Br	-5.4	+3.4	+2.2	-1.0
I	-32.2	+9.9	+2.6	-7.3
OH	+26.6	-12.7	+1.6	-7.3
OCH ₃	+31.4	-14.4	+1.0	-7.7
CHO	+8.2	+1.2	+0.6	+5.8
COCH ₃	+7.8	-0.4	-0.4	+2.8
COOH	+2.9	+1.3	+0.4	+4.3
CN	-16.0	+3.6	+0.6	+4.3
NO ₂	+19.6	-5.3	+0.9	+6.0
NH ₂	+19.2	-12.4	+1.3	-9.5
NHCOCH ₃	+11.1	-9.9	+0.2	-5.6
COOCH ₃	+2.0	+1.2	-0.1	+4.8