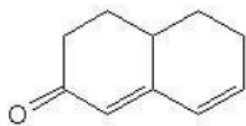




19. What increment will you add to the parent value for the exocyclic double bond marked in the structure?



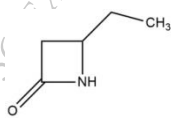
20. Draw the positive fragment generated by Retro Diel's Alder fragmentation of cyclohexene.

II. Long answer questions- (Answer any two)

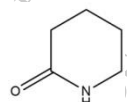
20M

1. a. A compound with molecular weight 137 has the following spectral characteristics  
 IR – 3358, 3080, 2868, 1598  $\text{cm}^{-1}$   
 $^1\text{H}$  NMR –  $\delta$  2.3 (bs, 2H, exchangeable), 2.5 (d, 2H), 4.25 (t, 1H),  
 5.3 (bs, 1H, exchangeable), 7.3 (s, 5H). Deduce structure and justify your answer.  
 Predict the  $m/z$  of its molecular ion peak.
1. b. How will you distinguish between the following compounds using suitable spectral techniques (i.e. UV, IR,  $^1\text{H}$  NMR,  $^{13}\text{C}$  NMR or MS). Give the differentiating spectra-

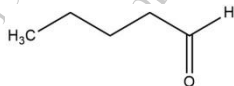
(i)



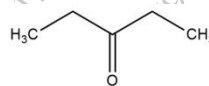
&



(ii)



&



2. a. Two isomers A and B with molecular formula of  $\text{C}_5\text{H}_{10}$  display the following  $^1\text{H}$  NMR spectra-

**Compound A:**  $\delta$  1.3(s, 18.3sq), 1.5(d, 8.9sq), 5.2(q, 3.2sq)

**Compound B:**  $\delta$  1.0(t, 4.8sq), 1.7(s, 5.1sq), 2.0(q, 3.1sq) 4.8(dd, 3.3sq)

Deduce the structures of A & B and justify your answer.

2. b. Write all possible structures for a compound with molecular formula  $\text{C}_4\text{H}_8\text{O}_2$  and whose IR spectrum shows a band at  $1740\text{ cm}^{-1}$  but does not show any band for O-H stretch. Predict  $^{13}\text{C}$  NMR for each

3. a. Elucidate the structure of compound with molecular formula  $\text{C}_{10}\text{H}_{11}\text{O}_2\text{Cl}$  and the following spectral details –

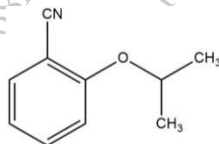
IR – 1745, 1600,  $1580\text{ cm}^{-1}$

$^1\text{H}$  NMR –  $\delta$  2.00(s, 3H), 2.8(t, 2H,  $J=6\text{ Hz}$ ), 4.1(t, 2H,  $J=6\text{ Hz}$ ), 7.1(d, 2H,  $J=8\text{ Hz}$ ), 7.3(d, 2H,  $J=8\text{ Hz}$ ).

Deduce the structure and justify your answer.

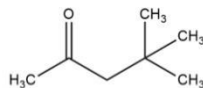


7. Predict: (i) IR and  $^{13}\text{C}$  NMR ( $\delta$  and splitting) of Compound M given below & (ii)  $^1\text{H}$  NMR spectrum of the Compound N given below giving chemical shifts, splitting pattern and integration.



M

&amp;



N

8. Compound M is an ether of molecular weight 136. It's MS data is as follows: m/z (intensity)-122(25), 95(6), 94(100), 77(8), 51(6), 43(5), 41(5), 39(6). Predict structure and explain the formation of base peak.
9. What is meant by supercritical fluid chromatography. Give 2 advantages of the technique over HPLC.

For Woodward Fieser Rule 11

## Conjugated dienes &amp; Trienes, Solvent: Ethanol

Parent value for Butadiene system or acyclic conjugated diene	217 nm
Acyclic triene	245 nm
Homoannular conjugated diene	253 nm
Heteroannular conjugated diene	215 nm
Increment for each substituents	
Alkyl substituents or ring residue	5 nm
Exocyclic double bond	5 nm
Double bond extending conjugation	30 nm
Auxochrome	
-OR	+6 nm
-SR	+30 nm
-Cl, -Br	+5 nm
NR <sub>2</sub>	+60 nm
-OCOCH <sub>3</sub>	0

Woodward Fieser rules for  $\alpha\beta$ -unsaturated carbonyl compounds:-

a) Parent values			
$\alpha\beta$ -unsaturated acyclic or six membered ketone	215 nm		
$\alpha\beta$ -unsaturated five membered ring ketone	202 nm		
$\alpha\beta$ -unsaturated aldehyde	207 nm		
b) Increments			
i) Each alkyl substituents or ring residue			
At $\alpha$ position	10 nm		
At $\beta$ position	12 nm		
At gamma and higher position	18 nm		
ii) Each exocyclic double bond	5 nm		
iii) Double bond extending conjugation	30 nm		
iv) Homoannular conjugated diene	39 nm		
Auxochromes	Positions		
	$\alpha$	$\beta$	gamma
-OH	35	30	50
-OR	35	30	17
-SR	-	85	-
-OCOCH <sub>3</sub>	6	6	6
-Cl	15	12	-
-Br	25	30	-
-NR <sub>2</sub>	-	95	-

IR Correlation Chart

## 3.4 Absorptions of Common Functional Groups

33

Table 3-1

CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS†

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
<b>A. Hydrocarbon chromophore</b>			
<b>1. C—H STRETCHING</b>			
a. Alkane	3.38–3.51	(m-s)	2962–2853
b. Alkene, monosubstituted (vinyl)	3.29–3.32	(m)	3040–3010
	and 3.23–3.25	(m)	3095–3075
Alkene, disubstituted, <i>cis</i>	3.29–3.32	(m)	3040–3010
Alkene, disubstituted, <i>trans</i>	3.29–3.32	(m)	3040–3010
Alkene, disubstituted, <i>gem</i>	3.23–3.25	(m)	3095–3075
Alkene, trisubstituted	3.29–3.32	(m)	3040–3010
c. Alkyne	~3.03	(s)	~3300
d. Aromatic	~3.30	(v)	~3030
<b>2. C—H BENDING</b>			
a. Alkane, C—H	~7.46	(w)	~1340
Alkane, —CH <sub>2</sub> —	6.74–6.92	(m)	1485–1445
Alkane, —CH <sub>3</sub>	6.80–7.00	(m)	1470–1430
	and 7.25–7.30	(s)	1380–1370
Alkane, <i>gem</i> -dimethyl	7.22–7.25	(s)	1385–1380
	and 7.30–7.33	(s)	1370–1365
Alkane, <i>tert</i> -butyl	7.17–7.22	(m)	1395–1385
	and ~7.33	(s)	~1365
b. Alkene, monosubstituted (vinyl)	10.05–10.15	(s)	995–985
	10.93–11.05	(s)	915–905
	and 7.04–7.09	(s)	1420–1410
Alkene, disubstituted, <i>cis</i>	~14.5	(s)	~690
Alkene, disubstituted, <i>trans</i>	10.31–10.42	(s)	970–960
	and 7.64–7.72	(m)	1310–1295
Alkene, disubstituted, <i>gem</i>	11.17–11.30	(s)	895–885
	and 7.04–7.09	(s)	1420–1410
Alkene, trisubstituted	11.90–12.66	(s)	840–790
c. Alkyne	~15.9	(s)	~630
d. Aromatic, substitution type: ‡			
five adjacent hydrogen atoms	~13.3	(v, s)	~750
	and ~14.3	(v, s)	~700
four adjacent hydrogen atoms	~13.3	(v, s)	~750
three adjacent hydrogen atoms	~12.8	(v, m)	~780
two adjacent hydrogen atoms	~12.0	(v, m)	~830
one hydrogen atom	~11.3	(v, w)	~880
<b>3. C—C MULTIPLE BOND STRETCHING</b>			
a. Alkene, nonconjugated	5.95–6.17	(v)	1680–1620
Alkene, monosubstituted (vinyl)	~6.08	(m)	~1645
Alkene, disubstituted, <i>cis</i>	~6.03	(m)	~1658
Alkene, disubstituted, <i>trans</i>	~5.97	(m)	~1675

† Abbreviations: s = strong, m = medium, w = weak, v = variable, b = broad, sh = sharp.  
~ = approximately

‡ Substituted benzenes also show weak bands in the region 5.0–6.0  $\mu$  (2000–1670  $\text{cm}^{-1}$ ) region that are characteristic of the substitution type. See Fig. 3-30.

## 34 INFRARED SPECTROSCOPY

## Chap. 3

Table 3-1 (cont.)

CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS†

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
Alkene, disubstituted, <i>gem</i>	~6.05	(m)	~1653
Alkene, trisubstituted	~5.99	(m)	~1669
Alkene, tetrasubstituted	~5.99	(w)	~1669
Diene	~6.06	(w)	~1650
and	~6.25	(w)	~1600
b. Alkyne, monosubstituted	4.67-4.76	(m)	2140-2100
Alkyne, disubstituted	4.42-4.57	(v, w)	2260-2190
c. Allene	~5.1	(m)	~1960
and	~9.4	(m)	~1060
d. Aromatic $\text{C}=\text{C}$	~6.25	(v)	~1600
	~6.33	(v)	~1580
	~6.67	(m)	~1500
and	~6.90	(m)	~1450
<b>B. Carbonyl chromophore</b>			
<b>I. KETONE STRETCHING VIBRATIONS</b>			
a. Saturated, acyclic	5.80-5.87	(s)	1725-1705
b. Saturated, cyclic:			
6-membered ring (and higher)	5.80-5.87	(s)	1725-1705
5-membered ring	5.71-5.75	(s)	1750-1740
4-membered ring	~5.63	(s)	~1775
c. $\alpha,\beta$ -Unsaturated, acyclic	5.94-6.01	(s)	1685-1665
d. $\alpha,\beta$ -Unsaturated, cyclic:			
6-membered ring (and higher)	5.94-6.01	(s)	1685-1665
5-membered ring	5.80-5.85	(s)	1725-1708
e. $\alpha,\beta,\alpha',\beta'$ -Unsaturated, acyclic	5.99-6.01	(s)	1670-1663
f. Aryl	5.88-5.95	(s)	1700-1680
g. Diaryl	5.99-6.02	(s)	1670-1660
h. $\alpha$ -Diketones	5.78-5.85	(s)	1730-1710
i. $\beta$ -Diketones (enolic)	6.10-6.50	(s)	1640-1540
j. 1,4-Quinones	5.92-6.02	(s)	1690-1660
k. Ketenes	~4.65	(s)	~2150
<b>2. ALDEHYDES</b>			
a. Carbonyl stretching vibrations			
Saturated, aliphatic	5.75-5.81	(s)	1740-1720
$\alpha,\beta$ -Unsaturated, aliphatic	5.87-5.95	(s)	1705-1680
$\alpha,\beta,\gamma,\delta$ -Unsaturated, aliphatic	5.95-6.02	(s)	1680-1660
Aryl	5.83-5.90	(s)	1715-1695
b. C-H Stretching vibrations, two bands	3.45-3.55	(w)	2900-2820
and	3.60-3.70	(w)	2775-2700
<b>3. ESTER STRETCHING VIBRATIONS</b>			
a. Saturated, acyclic	5.71-5.76	(s)	1750-1735
b. Saturated, cyclic:			
$\delta$ -lactones (and larger rings)	5.71-5.76	(s)	1750-1735
$\gamma$ -lactones	5.62-5.68	(s)	1780-1760
$\beta$ -lactones	~5.5	(s)	~1820

† Abbreviations: s = strong, m = medium, w = weak, v = variable, b = broad, sh = sharp.  
~ = approximately

## 3.4 Absor

Table 3-1 (CHARACTER

c. U

vi

a.

a.

a.

 $\beta$ .d.  $\alpha$ .e.  $\beta$ .

f. C

4. CAR

a. C

sa

a.

ar

b. H

se

c. C

5. ANH

VIBR

a. Sa

b.  $\alpha/\beta$ 

c. Sa

d.  $\alpha/\beta$ 

6. ACYL

VIBR

a. Ac

b. Ac

c. Ac

d.  $\alpha,\beta$ 

e. CC

f. CC

g. CO

7. AMIL

a. Ca

Pri

s

Pri

Sec

s

Sec

Ter

Cyc

Cyc

## 3.4 Absorptions of Common Functional Groups

35

Table 3-1 (cont.)

## CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
c. Unsaturated:			
vinyl ester type	5.56-5.65	(s)	1800-1770
$\alpha,\beta$ -unsaturated and aryl	5.78-5.82	(s)	1730-1717
$\alpha,\beta$ -unsaturated $\delta$ -lactone	5.78-5.82	(s)	1730-1717
$\alpha,\beta$ -unsaturated $\gamma$ -lactone	5.68-5.75	(s)	1760-1740
$\beta,\gamma$ -unsaturated $\gamma$ -lactone	~5.56	(s)	~1800
d. $\alpha$ -Ketoesters	5.70-5.75	(s)	1755-1740
e. $\beta$ -Ketoesters (enolic)	~6.06	(s)	~1650
f. Carbonates	5.62-5.75	(s)	1786-1740
4. CARBOXYLIC ACIDS			
a. Carbonyl stretching vibrations:			
saturated aliphatic	5.80-5.88	(s)	1725-1700
$\alpha,\beta$ -unsaturated aliphatic	5.83-5.92	(s)	1715-1690
aryl	5.88-5.95	(s)	1700-1680
b. Hydroxyl stretching (bonded), several bands			
	3.70-4.00	(w)	2700-2500
c. Carboxylate anion stretching			
	6.21-6.45	(s)	1610-1550
	and 7.15-7.69	(s)	1400-1300
5. ANHYDRIDE STRETCHING VIBRATIONS			
a. Saturated, acyclic			
	5.41-5.56	(s)	1850-1800
	and 5.59-5.75	(s)	1790-1740
b. $\alpha,\beta$ -Unsaturated and aryl, acyclic			
	5.47-5.62	(s)	1830-1780
	and 5.65-5.81	(s)	1770-1720
c. Saturated, 5-membered ring			
	5.35-5.49	(s)	1870-1820
	and 5.56-5.71	(s)	1800-1750
d. $\alpha,\beta$ -Unsaturated, 5-membered ring			
	5.41-5.56	(s)	1850-1800
	and 5.47-5.62	(s)	1830-1780
6. ACYL HALIDE STRETCHING VIBRATIONS			
a. Acyl fluorides			
	~5.41	(s)	~1850
b. Acyl chlorides			
	~5.57	(s)	~1795
c. Acyl bromides			
	~5.53	(s)	~1810
d. $\alpha,\beta$ -Unsaturated and aryl			
	5.61-5.72	(s)	1780-1750
	and 5.72-5.82	(m)	1750-1720
e. $\text{COF}_2$			
	5.19	(s)	1928
f. $\text{COCl}_2$			
	5.47	(s)	1828
g. $\text{COBr}_2$			
	5.47	(s)	1828
7. AMIDES			
a. Carbonyl stretching vibrations			
Primary, solid and concentrated solution	~6.06	(s)	~1650
Primary, dilute solution	~5.92	(s)	~1690
Secondary, solid and concentrated solution	5.95-6.14	(s)	1680-1630
Secondary, dilute solution	5.88-5.99	(s)	1700-1670
Tertiary, solid and all solutions	5.99-6.14	(s)	1670-1630
Cyclic, $\delta$ -lactams, dilute solution	~5.95	(s)	~1680
Cyclic, $\gamma$ -lactams, dilute solution	~5.88	(s)	~1700

Table 3-1 (cont.)  
CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS†

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
Cyclic, $\gamma$ -lactams, fused to another ring, dilute solution	5.71-5.88	(s)	1750-1700
Cyclic, $\beta$ -lactams, dilute solution	5.68-5.78	(s)	1760-1730
Cyclic, $\beta$ -lactams, fused to another ring, dilute solution	5.62-5.65	(s)	1780-1770
Ureas, acyclic	~6.02	(s)	~1660
Ureas, cyclic, 6-membered ring	~6.10	(s)	~1640
Ureas, cyclic, 5-membered ring	~5.81	(s)	~1720
Urethanes	5.75-5.92	(s)	1740-1690
Imides, acyclic	~5.85	(s)	~1710
and	~5.88	(s)	~1700
Imides, cyclic, 6-membered ring	~5.85	(s)	~1710
and	~5.88	(s)	~1700
Imides, cyclic, $\alpha,\beta$ -unsaturated, 6-membered ring	~5.78	(s)	~1730
and	~5.99	(s)	~1670
Imides, cyclic, 5-membered ring	~5.65	(s)	~1770
and	~5.88	(s)	~1700
Imides, cyclic, $\alpha,\beta$ -unsaturated, 5-membered ring	~5.59	(s)	~1790
and	~5.85	(s)	~1710
b. N-H Stretching vibrations			
Primary, free; two bands	~2.86	(m)	~3500
and	~2.94	(m)	~3400
Primary, bonded; two bands	~2.99	(m)	~3350
and	~3.15	(m)	~3180
Secondary, free; one band	~2.92	(m)	~3430
Secondary, bonded; one band	3.0-3.2	(m)	3320-3140
c. N-H Bending vibrations			
Primary amides, dilute solution	6.17-6.29	(s)	1620-1590
Secondary amides, dilute solution	6.45-6.62	(s)	1550-1510
C. Miscellaneous chromophoric groups			
I. ALCOHOLS AND PHENOLS			
a. O-H Stretching vibrations			
Free O-H	2.74-2.79	(v, sh)	3650-3590
Intermolecularly hydrogen bonded (change on dilution)			
single bridge compounds	2.82-2.90	(v, sh)	3550-3450
polymeric association	2.94-3.13	(s, b)	3400-3200
Intramolecularly hydrogen bonded (no change on dilution)			
single bridge compounds	2.80-2.90	(v, sh)	3570-3450
chelate compounds	3.1-4.0	(w, b)	3200-2500

† Abbreviations: s = strong, m = medium, w = weak, v = variable, b = broad, sh = sharp, ~ = approximately

## 3.4 Absorpti

Table 3-1 (con  
CHARACTERIST

CHARACTERIST

b. O—

vi

Prim

Seco

Tert

Phen

2. AMINI

a. N—

Prim

Seco

Imin

Amin

b. N—

Prim

Seco

Amin

c. C—

Aron

Aron

Aron

Aliph

3. UNSAT

COMPC

a. C≡N

Alkyl

$\alpha,\beta$ -U

Aryl

Isocy

Isocy

b. C=

(imin

Alkyl

$\alpha,\beta$ -U

c. —N=

azo

azo

d. —N=

diimic

e. —N<sub>3</sub>

## 38 INFRARED SPECTROSCOPY

## Chap. 3

Table 3-1 (cont.)

CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS†

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
f. C—NO <sub>2</sub> , Nitro compounds:			
aromatic	6.37-6.67	(s)	1570-1500
aliphatic	and 7.30-7.70	(s)	1370-1300
	6.37-6.45	(s)	1570-1550
	and 7.25-7.30	(s)	1380-1370
g. O—NO <sub>2</sub> , Nitrates	6.06-6.25	(s)	1650-1600
	and 7.70-8.00	(s)	1300-1250
h. C—NO, Nitroso compounds	6.25-6.67	(s)	1600-1500
i. O—NO, Nitrites	5.95-6.06	(s)	1680-1650
	and 6.15-6.21	(s)	1625-1610
4. HALOGEN COMPOUNDS, C—X, STRETCHING VIBRATIONS			
a. C—F	7.1-10.0	(s)	1400-1000
b. C—Cl	12.5-16.6	(s)	800-600
c. C—Br	16.6-20.0	(s)	600-500
d. C—I	~20	(s)	~500
5. SULFUR COMPOUNDS			
a. S—H Stretching vibrations	3.85-3.92	(w)	2600-2550
b. C=S Stretching vibrations	8.33-9.52	(s)	1200-1050
c. S=O Stretching vibrations:			
sulfoxides	9.35-9.71	(s)	1070-1030
sulfones	8.62-8.77	(s)	1160-1140
	and 7.41-7.69	(s)	1350-1300
sulfites	8.13-8.70	(s)	1230-1150
	and 7.00-7.41	(s)	1430-1350
sulfonyl chlorides	8.44-8.59	(s)	1185-1165
	and 7.30-7.46	(s)	1370-1340
sulfonamides	8.48-8.77	(s)	1180-1140
	and 7.41-7.69	(s)	1350-1300
sulfonic acids	8.27-8.70	(s)	1210-1150
	9.43-9.71	(s)	1060-1030
	and ~15.4	(s)	~650

† Abbreviations: s = strong, m = medium, w = weak, v = variable, b = broad, sh = sharp, ~ = approximately

3.4 Abs

Fig. 3-9 D

~1380  $\text{cm}^{-1}$  bending  
Simpl  
by an oxy  
C—O stre  
spectrum  
hydrocart  
a very pr  
C—O stre  
groups pr  
Fig. 3-10,  
is present  
polymeric  
(~1053  $\text{cm}^{-1}$ )  
alcohol. T

absorptions in terms of wavelengths or wave numbers. The remainder of this section is devoted to an examination of a number of infrared spectra. Significant structural features and absorptions are indicated.

The infrared spectrum of nujol (Fig. 3-3), a mixture of saturated hydrocarbons, contains absorptions resulting from vibrations typical of groups that are present in such molecules, i.e., C—H stretching (~3.39 and ~3.54  $\mu$ , ~2950 and ~2820  $\text{cm}^{-1}$ ), —CH<sub>2</sub>— bending (~6.86  $\mu$ , ~1458  $\text{cm}^{-1}$ ), and C—CH<sub>3</sub> bending (~6.86 and 7.28  $\mu$ , ~1458 and

Fig. 3-10 Lairy

## 3.4 Absorptions of Common Functional Groups

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Table 3-1 (cont.)

## CHARACTERISTIC INFRARED ABSORPTIONS OF FUNCTIONAL GROUPS

Group	Range $\mu$	Intensity	Range $\text{cm}^{-1}$
<b>b. O—H Bending and C—O stretching vibrations</b>			
Primary alcohols	$\sim 9.5$	(s)	$\sim 1050$
	and 7.4-7.9	(s)	1350-1260
Secondary alcohols	$\sim 9.1$	(s)	$\sim 1100$
	and 7.4-7.9	(s)	1350-1260
Tertiary alcohols	$\sim 8.7$	(s)	$\sim 1150$
	and 7.1-7.6	(s)	1410-1310
Phenols	$\sim 8.3$	(s)	$\sim 1200$
	and 7.1-7.6	(s)	1410-1310
<b>2. AMINES</b>			
<b>a. N—H Stretching vibrations</b>			
* Primary, free; two bands	$\sim 2.86$	(m)	$\sim 3500$
	and $\sim 2.94$	(m)	$\sim 3400$
Secondary, free; one band	2.86-3.02	(m)	3500-3310
Imines (=N—H); one band	2.94-3.03	(m)	3400-3300
Amine salts	3.2-3.3	(m)	3130-3030
<b>b. N—H Bending vibrations</b>			
Primary	6.06-6.29	(s-m)	1650-1590
Secondary	6.06-6.45	(w)	1650-1550
Amine salts	6.25-6.35	(s)	1600-1575
	and $\sim 6.67$	(s)	$\sim 1500$
<b>c. C—N Vibrations</b>			
Aromatic, primary	7.46-8.00	(s)	1340-1250
Aromatic, secondary	7.41-7.81	(s)	1350-1280
Aromatic, tertiary	7.36-7.64	(s)	1360-1310
Aliphatic	8.2-9.8	(w)	1220-1020
	and $\sim 7.1$	(w)	$\sim 1410$
<b>3. UNSATURATED NITROGEN COMPOUNDS</b>			
<b>a. C<math>\equiv</math>N Stretching vibrations</b>			
Alkyl nitriles	4.42-4.46	(m)	2260-2240
$\alpha, \beta$ -Unsaturated alkyl nitriles	4.47-4.51	(m)	2235-2215
Aryl nitriles	4.46-4.50	(m)	2240-2220
Isocyanates	4.40-4.46	(m)	2275-2240
Isocyanides	4.50-4.83	(m)	2220-2070
<b>b. C=N— Stretching vibrations (imines, oximes)</b>			
Alkyl compounds	5.92-6.10	(v)	1690-1640
$\alpha, \beta$ -Unsaturated compounds	6.02-6.14	(v)	1660-1630
<b>c. —N=N— Stretching vibrations, azo compounds</b>			
	6.14-6.35	(v)	1630-1575
<b>d. —N=C=N— Stretching vibrations, diimides</b>			
	4.64-4.70	(s)	2155-2130
<b>e. —N<sub>3</sub> Stretching vibrations, azides</b>			
	4.63-4.72	(s)	2160-2120
	and 7.46-8.48	(w)	1340-1180



Proton Chemical Shifts (Values are given on the officially approved  $\delta$  scale;  $\tau = 10.00 - \delta$ )

Substituent Group	Methyl Protons	Methylene Protons	Methine Proton
HC-C-CH <sub>3</sub>	0.95	1.20	1.55
HC-C-NR <sub>2</sub>	1.05	1.45	1.70
HC-C-C=C	1.00	1.35	1.70
HC-C-C=O	1.05	1.55	1.95
HC-C-NR <sub>2</sub> Ar	1.10	1.50	1.80
HC-C-NH(C=O)R	1.10	1.50	1.90
HC-C-(C=O)NR <sub>2</sub>	1.10	1.50	1.80
HC-C-(C=O)Ar	1.15	1.55	1.90
HC-C-(C=O)OR	1.15	1.70	1.90
HC-C-Ar	1.15	1.55	1.80
HC-C-OH (and OR)	1.20	1.50	1.75
HC-C-C≡C-R	1.20	1.50	1.80
HC-C-C≡N	1.25	1.65	2.00
HC-C-SR	1.25	1.60	1.90
HC-C-OAr	1.30	1.55	2.00
HC-C-O(C=O)R	1.30	1.60	1.80
HC-C-SH	1.30	1.60	1.80
HC-C-(S=O)R and -SO <sub>2</sub> R	1.35	1.60	1.65
HC-C-NR <sub>3</sub>	1.40	1.75	2.05
HC-C-O(C=O)CF <sub>3</sub>	1.40	1.65	1.95
HC-C-Cl	1.55	1.80	1.85
HC-C-O(C=O)Ar	1.65	1.75	1.90
HC-C-Br	1.80	1.85	1.90
HC-CH <sub>2</sub>	0.90	1.30	1.50
HC-C=C	1.60	2.05	2.80
HC-C≡C	1.70	2.20	2.50
HC-(C=O)OR (and NR <sub>3</sub> )	2.00	2.25	3.00
HC-SR	2.05	2.55	3.00
HC-O-O	2.10	2.30	2.55
HC-(C=O)R	2.10	2.35	2.65
HC-C≡N	2.15	2.45	2.90
HC-CHO	2.20	2.40	2.85
HC-Ar (and NR <sub>2</sub> )	2.25	2.45	3.40
HC-SSR	2.35	2.70	3.60
HC-(C=O)Ar	2.40	2.70	3.60
HC-SAr	2.40	2.70	3.60
HC-NR <sub>2</sub> Ar	2.60	3.10	3.60
HC-SO <sub>2</sub> R and -(SO)R	2.60	3.05	4.10
HC-Br	2.70	3.40	3.60
HC-NR <sub>3</sub>	2.95	3.10	3.85
HC-NH(C=O)R	2.95	3.35	4.05
HC-Cl	3.05	3.45	3.60
HC-OH and -OR	3.20	3.40	4.05
HC-NH <sub>2</sub>	3.50	3.75	4.05
HC-O(C=O)R	3.65	4.10	4.95
HC-OAr	3.80	4.00	4.60
HC-O(C=O)Ar	3.80	4.20	5.05

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Substituent Group	Methyl Protons	Methylene Protons	Methine Proton
HC-F	4.25	4.50	4.80
HC-NO <sub>2</sub>	4.30	4.35	4.60
Cyclopropane	0.20	2.45	0.40
Cyclobutane	1.65	1.50	1.80
Cyclopentane	1.50	1.25	
Cyclohexane	1.50	1.25	
Cycloheptane	1.25	1.25	


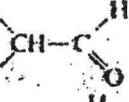
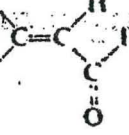





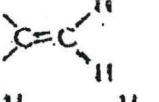



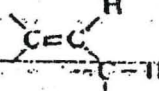
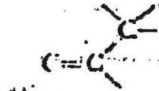

Substituent Group	Proton Shift	Substituent Group	Proton Shift
HC≡CH	2.35	HO-C=O	10-12
HC≡CAr	2.90	HO-SO <sub>2</sub>	11-12
HC≡C-C=C	2.75	HO-Ar	4.5-6.5
HA-Ar	7.20	HO-R	0.5-4.5
HCO-O	8.1	HS-Ar	2.8-3.6
HCO-R	9.4-10.0	HS-R	1-2
HCO-Ar	9.7-10.5	HN-Ar	3-6
HO-N=C(oxime)	9-12	HN-R	0.5-5

<sup>13</sup>C Chemical Shifts (Values given on the  $\delta$  scale, relative to TMS.)

Substituent Group	Primary Carbon	Secondary Carbon	Tertiary Carbon	Quaternary Carbon
Alkanes	-20 to 30	25 to 45	30 to 60	35 to 70
C-C=C	40 to 60	40 to 70	60 to 75	70 to 85
C-O	20 to 45	40 to 60	50 to 70	65 to 75
C-N	10 to 30	25 to 45	40 to 55	55 to 70
C-S	-37 to 35	-10 to 45	30 to 65	35 to 75
C-Halide	(I), (Cl)	(I), (Cl)	(I), (Cl)	(I), (Cl)
Alkenes	70 to 100	130 to 150	130 to 150	130 to 150
Alkynes	110 to 150	110 to 135	110 to 135	150 to 160
Aromatics	125 to 145	115 to 140	115 to 140	155 to 165
C-substituted Heteroaromatics	135 to 155	105 to 120	105 to 120	150 to 170
C- $\alpha$	115 to 140	115 to 140	115 to 140	165 to 185
Cyanates	115 to 135	115 to 135	115 to 135	150 to 175
Isocyanates	110 to 120	110 to 120	110 to 120	160 to 180
Thiocyanates	110 to 120	110 to 120	110 to 120	160 to 185
Isothiocyanates	120 to 140	120 to 140	120 to 140	175 to 205
Cyanides	110 to 130	110 to 130	110 to 130	175 to 225

TURN OVER

## SPIN-SPIN COUPLING CONSTANTS

Type	J, cps	Type	J, cps
$H_2^+$	280	$C-CH=CH-C$	9-13
$CH_2^+$	12-4	$H-C\equiv C-H^+$	9-1
	12-15	$CH-C\equiv C-H$	2-3
$CH-CH$	2-9		1-3
$-C(-H)-C(-H)-C(-H)-$	-0		6-8
$CH_2-CH_2-X$	6.5-7.5		<i>o</i> -6-9 <i>m</i> -1-3 <i>p</i> -0-1
$CH_2-CH_2-N$	5.5-7.0		<i>ortho</i> 1.6-2.0 <i>meta</i> 0.6-1.0 <i>para</i> 1.3-1.8 <i>ortho</i> 3.2-3.8
$CH_2$			<i>ortho</i> 2.0-2.6 <i>meta</i> 1.5-2.2 <i>para</i> 1.8-2.3 <i>ortho</i> 2.8-4.0
	<i>ortho</i> 5-10 <i>meta</i> 2-4 <i>cis</i> 2-4		<i>ortho</i> 4.6-5.8 <i>meta</i> 1.0-1.8 <i>para</i> 2.1-3.3 <i>ortho</i> 5.0-4.2
	0.5-3		<i>ortho</i> 4.9-5.7 <i>meta</i> 1.6-2.6
	7-12		<i>ortho</i> 0.7-1.1 <i>meta</i> 0.2-0.5 <i>ortho</i> 7.2-8.5 <i>ortho</i> 1.4-1.9
	13-18		
	4-10		
	<i>ortho</i> 0.5-2.5		
	-0		

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